(12)

EUROPEAN PATENT APPLICATION published in accordance with Art. 158(3) EPC

(43) Date of publication: 06.06.2001 Bulletin 2001/23

(21) Application number: 99937053.9

(22) Date of filing: 12.08.1999

(51) Int Cl.7: **C07D 471/04**, C07D 471/14, C07D 491/113, C07D 495/14, A61K 31/435, A61K 31/47

(86) International application number: PCT/JP99/04381

(87) International publication number: WO 00/09506 (24.02.2000 Gazette 2000/08)

(84) Designated Contracting States: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE Designated Extension States:

(30) Priority: 12.08.1998 JP 24106298 30.07.1999 JP 21612599

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(54) 1H-IMIDAZOPYRIDINE DERIVATIVES

(57) 1H-imidazopyridine derivatives represented by the following general formula or salts thereof:

wherein R¹ represents hydrogen atom, hydroxyl group, an alkyl group, a cycloalkyl group, styryl group, or an aryl group; R² represents hydrogen atom, an alkyl group, a halogen atom, hydroxyl group, amino group, cyclic amino group, or phenoxy group; ring A represents a homocyclic or heterocyclic ring which may be substituted; R² represents a saturated hitrogen-containing heterocyclic group; and m represents an integer of from 0 to 3. The derivatives have excellent inhibitory actions against production of TNF or IL-1 and are actiremely useful as preventive or therapeutic agents for diseases in which a cyclokin is mediated.

Description

Technical Field

9 (0001) The present invention relates to novel 1H-imidazopyridine derivatives or salts thereof which have a potent inhibitory action against production of tumor necrotizing factor (TNF) or interleukin-1 (IL-1) and are useful as medicaments for preventive or therapeutic treatment of diseases of humans and animals, in which a cytokine such as TNF, IL-1 is mediated, which include chronic inflammatory diseases (e.g., reumatic arthritis, osteoarthritis, etc.), allergic rhillitis, aotopic dermatilis, contact dermatilis, as stamba, spesis, sopici shock, various autiorimune diseases (act.), allergic rhillitis, actions autionimune diseases (act.), antiperior diseases (e.g., hemolytic anemia, anaplastic anemia, idiopathic thrombocythemia, etc.), autoimmune intestinal diseases (e.g., ubcrative colitis, Crohn's diseases, etc.), autoimmune comettis (e.g., keratoconjunctivitis sicca, spring calarm, etc.), endocrine ophthalmopathy. Graves disease, sarcold granuloma, multiple scherosis, systemic erythematodes, multiple chordrifis, pachydermia, active chronic hepatitis, myasthenia gravis, psoriasis, interstitial pulmonary tibrosis and the like.

Background Art

[0002] Some compounds having 1H-imidazoquinoline structure are known which are analogous to the compounds of the present invention. Journal of Medicinal Chemistry, Vol. 11, p. 87 (1989) discloses 1-(2-piperidinocity)-1H-imidazo(4.5-0)-quinoline, Japanese Patent Unexamined Publication (KOKAI) No. 5ho 60-122488/1985 discloses 1-isouty:1-H-imidazo(4.5-0)-quinoline-4-amine (general name: imiquimod) as a compound having an antiviral action, and Hungarian Patent Publication No. 3479 (Patent No. 190109) discloses 1-(2-distriparimicothy)-1H-imidazo(4.5-0)-quinoline as a compound having analogesic and anticonvulsant actions. However, 1H-imidazopyridine derivatives as those according to the present invention have never been known so far.

[0003] Moreover, the aforementioned imiquimod has been known to have an inducing action of a few kinds of cytokines such as interferon (IFN), TNF, IL-1 and the like, which is described in Journal of Interferon Research, Vol. 14, p. 81 (1994). However, 1H-Imidazopyridine derivatives or 1H-Imidazoquinoline derivatives having an inhibitory action against production of TNF or IL-1, which action is totally opposite to those taught by the aforementioned prior arts, have never been knowns of an

Disclosure of the Invention

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[0004] An object of the present invention is to provide novel compounds which have excellent inhibitory actions against production of cytokines such as TNF and IL-1 and the like are useful as medicaments.

[0005] The inventors of the present invention made intensive studies to achieve the object. As a result, they found novel 1H-imidazopyridine derivative which have an excellent inhibitory action against production of TNF or IL-1 and achieved the present invention.

[0006] The present invention thus relates to novel 1H-imidazopyridine derivatives represented by the following general formula (I) or salts thereof:

wherein R¹ represents hydrogen atom, hydroxyl group, an alkyl group which may have one or more substituents, a cycloalkyl group which may be substituted, a stry group which may be substituted, or an any group which may he one or more substituents. Pre-greensents hydrogen atom, an alkyl group, a halogen atom, hydroxyl group, an amino group which may have one or two substitutents, a cyclic amino group which may be substituted, or a phenoxy group which may be substituted, fing A represents a homocyclic or heterocyclic ring which may be substituted with one or more alkyl groups, alkoxyl groups, or halogen atoms; R³ represents a saturated nitrogen-containing heterocyclic group which may be substituted; and m represents an integer of from 0 to 2; provided that, when R³ represents unsubstituted piperdidno group, at least one of R³ and R² is not hydrogen atom.

[0007] According to the second embodiment of the present invention, there are provided novel 1H-imidazopyridine

derivatives represented by the following general formula (II) or salts thereof:

wherein R1, R2, ring A andm have the same meanings as those defined above; R4 represents hydrogen atom, an alkyl group, benzyl group, triphenylmethyl group, an alkanoyl group which may be substituted, an alkanosurfornyl group, athiocarbamoyl group which may be substituted, an alkanesulfonyl group, a benz renesulfonyl group, which may be substituted, or amiding group; Y represents methylene group, oxygen atom, sulfur atom, nitrogen atom, a croup represented to Whi. Or a single book; and ne prosessent an integer of from 0 to 2.

[9008] According to the third embodiment of the present invention, there are provided, among the compounds represented by the aforementioned general formulas (i) and (ii), the compounds wherein ring A is a benzene ring or a thiothera ring or the selfs thereof

thiophene ring, or the salts thereof. [0009] According to another aspect, there is provided a medicament which comprises as an active ingredient the compound represented by the aforementioned general formula (I) or (II), or a pharmacologically acceptable sait thereof. The medicament is useful for preventive or therapeutic treatment of diseases of mammals including humans, in which a cytokine such as TNF, IL-1 is mediated, which include chronic inflammatory diseases (e.g., rheumatic arthritis, osteoarthritis, etc.), allergic rhinitis, atopic dermatitis, contact dermatitis, asthma, sepsis, septic shock, various autoimmune diseases (autoimmune hemic diseases (e.g., hemolytic anemia, anapiastic anemia, idiopathic thrombocythemia, etc.), autoimmune intestinal diseases (e.g., ulcerative colitis, Crohn's disease, etc.), autoimmune comeitis (e.g., keratoconjunctivitis sicca, spring catarrh, etc.), endocrine ophthalmopathy, Graves disease, sarcoid granuloma, multiple sclerosis, systemic erythematodes, multiple chondritis, pachydermia, active chronic hepatitis, myasthenia gravis, psoriasis, interstitial pulmonary fibrosis and the like], diabetes, cancerous cachexia, HIV-infectious cachexia and the like. [0010] According to a further aspect, there are provided a use of the compound represented by the aforementioned general formula (I) or (II), or a pharmacologically acceptable salt thereof for the manufacture of the aforementioned medicament; and a method for the preventive or therapeutic treatment of diseases in which a cytokine such as TNF, IL-1 is mediated, which comprises the step of administering a preventively or therapeutically effective amount of the compound represented by the aforementioned general formula (f) or (ff), or a pharmacologically acceptable salt thereof to a mammal including a human. In addition, the present invention provides an inhibitor against production of tumor necrotizing factor (TNF) or interleukin-1 (IL-1) which comprises as an active ingredient the compound represented by the aforementioned general formula (I) or (II), or a pharmacologically acceptable salt thereof.

49 Best Mode for Carrying Out the Invention

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[0011] Specific explanations of the compounds of the aforementioned general formulas (i) and (ii) of the present invention will be given below. The compounds represented by the aforementioned general formula (ii) are characterized in that they have a specific saturated nitrogen-containing heterocyclic group which may have specific substituents as R3 among the compounds represented by the aforementioned general formula (ii). However, the scope of the present invention is not limited to the compounds represented by the aforementioned general formula (ii), and it should be understood that any compounds having as R3 a saturated nitrogen-containing heterocyclic group which may be substituted fall within the scope of the present invention.

[0012] In the aforementioned general formulas (I) and (II), examples of the alkyl group represented by R¹, R² or R⁴ include, for example, methyl group, eithyl group, propoly group, is propryl group, probuly group, isobulyl group, sec-bulyl group, pre-pentyl group, isopentyl group, neopentyl group, nebyl group and the like.

[0013] Examples of the cycloality group represented by R1 include, for example, cycloproxy group, cyclobuty if group, cycloproxy group, and include, for example, phenyl group, 2-pyridyl group, 3-pyridyl group, 3-pyridyl group, 3-pyridyl group, 3-pyridyl group, 3-pyridyl group, 2-pyridyl group, 2-pyridyl group, 2-pyridyl group, 2-pyridyl group, 2-pyridyl group, 2-pyridyl group, 3-pyridyl group, 1-imidazolyl group, 2-imidazolyl group, 2-pyrazolyl group, 3-pyrazolyl group, 3

azoly group, 3-isothiazolyl group, 4-isothiazoly group, 5-isothiazolyl group, 1,2-3-triazol-1y group, 1,2-3-triazol-1y group, 1,2-4-triazol-1y group, 1,2-4-triazol-1y group, 1,2-4-triazol-1y group, 1-4-triazol-1y group, 5-isotrazolyl group, 1-2-5-thiadiazol-3y group, 1-indolyl group, 2-indolyl group, 3-indolyl group and the like. [0014] Examples of the halogen atom represented by R8 include, for example, fluorine atom, chlorine atom, chlorine atom, and iodine atom. Examples of the amino group which may have one or two substituents that is represented by R8 include, for example, amino group, mothlyamino group, pythamino group, repropylamino group, isopropylamino group, cyclobrylamino group, cyclopropylamino group, group, group, group, group, group, dimethylamino group, and group, gro

group, cyclopropylamino group, cyclobulylamino group, cyclopentylamino group, cyclopentylamino group, dimethylamino group, barylamino group, barylamino group, barylamino group, barylamino group, barylamino group, paralmino group paralmino group paralmino group paralmino group paralmino group paralmino group, 1-paralmino group, 1-paral

[0015] Examples of the homocyclic or heterocyclic ring represented by ring A in the aforementioned general formulas (f) and (ii) include, for example, benzene ring, cyclopentene ring, cyclobentene ring, cyclobeptane ring, production ring, cyclobeptane ring, britisher ring, britisher ring, production ring, cyclobeptane ring, britisher ring, britisher ring, production ring, cyclobeptane ring, and the like. Examples of the alkyl group which may be substituted on the homocyclic or heterocyclic ring include, for example, methyl group, aboutly group, properly group, sepropyl group, n-butyl group, sibotyl group, sebotyl group, produced produced ring included, for example, methoxy group, althory group, n-propoxy group, isopropoxy group, n-butoxy group, isopropoxy group, isopropoxy group, n-butoxy group, isobutoxy group, and the like. Examples of the alkoyd group, seboty group, isopropoxy group, isopropoxy group, n-butoxy group, isobutoxy group, and the like. Examples of the alkogen atom which may be substituted on the said ring include, for example, fluorine atom, chlorine atom, promine atom, and addine atom. The number and kind of these substituents are not particularly limited, and when two or more substituents exist, they may be the same or different.

[0016] In the aforementioned general formula (I), the saturated nitrogen-containing heterocyclic group represented by RP means a saturated nitrogen-containing heterocyclic group which has one or more nitrogen atoms as ring-constituting atoms. Brampies include 1-aziridinyl group, 2-aziridinyl group, 2-aziridinyl group, 3-aziridinyl group, 3-aziridinyl group, 3-prollidinyl group, 3-aziridinyl group, 3-prollidinyl group, 3-prollidinyl group, 3-prollidinyl group, 3-prollidinyl group, 3-prollidinyl group, 4-properatinyl group, 3-prollidinyl group, 4-properatinyl group, 3-prollidinyl group, 4-prollidinyl group, 3-prollidinyl group, 4-prollidinyl group, 3-prollidinyl group, 4-prollidinyl group, 5-prollidinyl group, 4-prollidinyl group, 6-prollidinyl group, 6-pro

group, 3-ezetidinyl group, 2-morpholinyl group, 2-thiomorpholinyl group and the like.

[0017] In the aforementioned general formula (II), exemples of the alkanoy group which may be substituted that is presented by H include, for example, formly group, actify group, propionyl group, nothyryl group, isourlayly group, pivaloyl group, illuoraacetyl group, citilioraacetyl group, properties of the alkoxycarbonyl gr

[0018] In the present specification, with respect to the substituting/binding position of the terms "the any group", "the homocyclic or heterocyclic ring" and "saturated nitrogen-containing heterocyclic group", the terms herein used encompass any groups in their meanings which may substitute/bind at any position on a substitutable/bondable element among ring-constituting atoms, so long as the substituting/binding position is not particularly limited, as some examples are shown above.

[0019] In the aforementioned general formulas (I) and (II) of the present invention, when certain functional groups are referred to as 'which may be substituted' or 'which may have substitutents," the substituent may be any group so long as it can substitute on the functional groups. The number and kind of the substituent are not particularly limited, and when two or more substituents exist. they may be the same or different. Examples include halopen atoms such

as fluorine atom, chlorine atom, and bromine atom; hydroxyl group; alkyl groups such as methyl group, ethyl group, n-propyl group, isopropyl group, n-butyl group, isobutyl group, sec-butyl group, tert-butyl group, n-pentyl group, isopentyl group, neopentyl group, and n-hexyl group; trifluoromethyl group; aryl groups such as phenyl group, naphthyl group, and pyridyl group; alkoxyl groups such as methoxy group, ethoxy group, n-propoxy group, isopropoxy group, n-butoxy group, isobutoxy group, sec-butoxy group, and tert-butoxy group; aryloxy groups such as phenoxy group; amino groups which may be substituted such as amino group, methylamino group, ethylamino group, n-propylamino group, isopropylamino group, cyclopropylamino group, cyclobutylamino group, cyclopentylamino group, cyclohexylamino group, dimethylamino group, diethylamino group, anilino group, pyridylamino group, benzylamino group, dibenzylamino group, acetylamino group, trifluoroacetylamino group, tert-butoxycarbonylamino group, benzyloxycarbonylamino group, benzhydrylamino group, and triphenylmethylamino group; formyl group; alkanoyl groups such as acetyl group, propionyl group, n-butyryl group, isobutyryl group, valeryl group, isovaleryl group, pivaloyl group, fluoroacetyl group, diffuoroacetyl group, trifluoroacetyl group, chloroacetyl group, dichloroacetyl group, and trichloroacetyl group; alkoxycarbonyl groups such as methoxycarbonyl group, ethoxycarbonyl group, n-propoxycarbonyl group, isopropoxycarbonyl group, n-butoxycarbonyl group, isobutoxycarbonyl group, sec-butoxycarbonyl group, tert-butoxycarbonyl group, n-pentyloxycarbonyl group, and n-hexyloxycarbonyl group; benzyloxycarbonyl group; carbamoyl group; alkylcarbamoyl groups such as methylcarbamoyl group, ethylcarbamoyl group, n-propylcarbamoyl group, isopropylcarbamoyl group, n-butylcarbamoyl group, isobutylcarbamoyl group, sec-butylcarbamoyl group, and tert-butylcarbamoyl group; thiocarbamoyl group; alkylthiocarbamoyl groups such as methylthiocarbamoyl group, ethylthiocarbamoyl group, n-propylthiocarbamoyl group, isopropylthiocarbamoyl group, n-butylthiocarbamoyl group, isobutylthiocarbamoyl group, sec-butylthiocarbamoyl group, and tert-butylthiocarbamoyl group; amidino group; alkylthio groups such as methylthio group; alkanesulfinyl groups such as methanesulfinyl group; alkanesulfonyl groups such as methanesulfonyl group, ethanesulfonyl group, n-propanesulfonyl group, and n-butanesulfonyl group; arylsulfonyl groups such as ptoluenesulfonyl group, p-methoxybenzenesulfonyl group, and p-fluorobenzenesulfonyl group; aralkyl groups such as benzyl group, naphthyl group, pyridylmethyl group, furfuryl group, and triphenylmethyl group; nitro group; cyano group; sulfamoyl group; oxo group; hydroxyimino group; alkoxyimino groups such as methoxyimino group, ethoxyimino group, n-propoxylmino group, and isopropoxylmino group; ethylenedioxy group and the like.

[0020] The compounds represented by the aforementioned general formulas (i) and (ii) of the present invention can be converted into salts, preferably, pharmacologically acceptable salts, if desired, or free bases can be generated from the resulting salts.

[0021] Exemples of the salts, preferably, the pharmacologically acceptable salts, of the compounds represented by the aforementioned general formulae (i) and (ii) of the present invention include acid-addition salts, for example, salts with mineral acids such as hydrochloric acid, hydrobromic acid, hydroiodic acid, nitric acid, sultric acid, and phosphoric acid, and salts with organic acids such as a cette acid, propionic acid, formic acid, formic acid, waleric acid, maleic acid, furnaric acid, citric acid, examic acid, suchic acid, all, acite acid, methanesulforic acid, chanesulforic acid, sometiments acid, such acid, acid, acid acid, acid acid, acid acid, acid, acid, acid, sometiments acid, stearic acid, gluconic acid, inclotinic acid, tilluropracetic acid, and benzole acid.

[0022] Among the compounds represented by the aforementioned general formulas (i) and (ii) of the present invention, optical isomers may exist for compounds having asymmetric carbons. These optical active compounds and mixtures thereof fall within the scope of the present invention.

© [0023] The compounds represented by the aforementioned general formulas (I) and (II) or the salts thereol according to the present invention can exist as any crystalline form depending on manufacturing conditions, or exist as any hydrate or solvate. These crystalline forms, hydrates or solvates, and mixtures thereof fall within the scope of the present invention.

[0024] Preferred compounds of the present invention include, for example, the following compounds and salts there-

- (1) 4-chloro-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (2) 4.8-dichloro-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (3) 4-chloro-8-methyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (4) 4-chloro-8-methoxy-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4.5-clquinoline:

- (5) 4-chloro-2-phenyl-1-[2 -(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (6) 4.8-dichloro-2-phenyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (7) 4-chloro-8-methyl-2-phenyl-1-[2-(4-piperidyl)ethyll-1H-imidazo[4.5-c]quinoline:
- (8) 4-chloro-8-methoxy-2-phenyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (9) 4-chloro-1-[2-(4-piperidyl)ethyl]-2-trifluoromethyl-1H-imidazo[4,5-c]quinoline;
 - (10) 4,8-dichloro-1-[2-(4-piperidyl)ethyl]-2 -trifluoromethyl-1H-imidazo[4,5-c]quinoline;
 - (11) 4-chloro-8-methyl-1-[2-(4-piperidyl)ethyl]-2-trifluoromethyl-1H-imidazo[4,5-clquinoline;
 - (12) 4-chloro-8-methoxy-1-[2-(4-piperidyl)ethyl]-2-trifluoromethyl-1H-imidazo[4,5-c]quinoline;

- (13) 4-chloro-2-(4-methylphenyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (14) 4-chloro-2-(4-methoxyphenyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-clquinoline:
- (15) 4-chloro-2-(4-fluorophenyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (16) 4-chloro-1-[2 -(4-piperidyl)ethyl]-2-(4-trifluoromethylphenyl)-1H-imidazo[4,5-c]quinoline;
- (10) 4-cnloro-1-[z -[4-piperlayi)ethyl]-2-[4-trilluoromethylpnenyi)-1H-imidazo[4,5-c]qt (17) 4-chloro-2-(2-furyl)-1-[2-(4-piperldyl)ethyl]-1H-imidazo[4,5-c]quinoline;
 - (18) 4-chloro-1-[2-(4-piperidyl)ethyl]-2-(2-thienyl)-1H-imidazo[4,5-c]quinoline;
 - (19) 4-chloro-2-(2-imidazolvl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-clquinoline:
 - (20) 4-chloro-1-[2-(4-piperidyl)ethyl]-2-(2-thiazolyl)-1H-imidazo[4,5-c]quinoline;
 - (21) 4-chloro-2-(5-methyl-2-thienyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
 - (22) 4-chloro-1-[2-(4-piperidyl)ethyl]-2-(2-pyrrolyl)-1H-imidazo[4,5-c]quinoline;

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- (22) 4-critoro-1-[2-(4-piperidyl)etryl]-2-(2-pyrrolyl)-1H-imidazo[4,5-c]quinoline;
- (24) 2-(4-fluorophenyl)-4-methyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-clquinoline;
- (25) 4-methyl-1-[2-(4-piperidyl)ethyl]-2-(4-trifluoromethylphenyl)-1H-imidazo[4,5-c]quinoline,
- (26) 2-(2-furyl)-4-methyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
- (27) 4-methyl-1-[2-(4-piperidyl)ethyl]-2-(2-thienyl)-1H-imidazo[4,5-c]quinoline;
 - (28) 2-(2-imidazolyl)-4-methyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
 - (29) 4-methyl-1-[2-(4-piperidyl)ethyl]-2-(2-thiazolyl)-1H-imidazo[4,5-c]quinoline;
 - (30) 4-methyl-2-(3-methyl-2-thienyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4.5-c]quinoline;
 - (31) 4-methyl-2-(5-methyl-2-thienyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4.5-c]quinoline;
 - (32) 4-methyl-1-[2-(4-piperidyl)ethyl]-2-(2-pyrrolyl)-1H-imidazo[4,5-c]quinoline;
 - (33) 4-methyl-2-(1-methyl-2-pyrrolyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
 - (34) 4-chloro-6,7,8,9-tetrahydro-2-phenyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline;
 - (35) 4-chloro-6,7-dihydro-2-phenyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[5,4-d]cyclopenta[b]pyridine;
 - (36) 4-chloro-2-phenyl-1-[2-(4-pipendyl)ethyl]-1H-imidazo[5,4-d]thieno-[3,2-b]pyridine;
- 25 (37) 4-chloro-2-phenyl-1-[2-(3-pipendyl)ethyl]-1H-imidazo[4,5-c]quinoline;
 - (38) 4-chloro-1-[2-(2-morpholinyl)ethyl]-2-phenyl-1H-imidazo[4,5-c]quinoline;
 - (39) 4-chloro-2-phenyl-1-[2-(1-piperazinyl)ethyl]-1H-imidazo[4,5-c]quinoline:
 - (40) 4,6,7,8,9-pentachloro-2-ethoxymethyl-1-[2-(4-thiomorpholinyl)ethyl]-1H-imidazo[4,5-c]quinoline;
 - (41) 4-chloro-6,7,8,9-tetrahydro-2-hydroxymethyl-1-[2-(1-piperazinyl)ethyl]-1H-imidazo[5,4-d]cyclohepta[b]pyrid-iner and
 - (42) 4-chloro-2-(3-methyl-2-thienyl)-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline.

[0025] The novel 1H-imidazopyridine derivatives represented by the aforementioned general formula (I) or (II) according to the present invention can be prepared by various methods; however, the preparation methods of the compounds of the present invention are not limited thereto. In the following preparation methods, specific explanations for the compounds represented by the aforementioned general formula (I) will be given, and it is obvious that these prepration methods include the comounds represented by the aforementioned general formula (II).

[0026] As the first synthetic method of the compounds of the present invention, the following synthetic method can be used in accordance with the method disclosed in Japanese Patent Unexamined Publication (KOKAI) No. Hel 3-20678/1991 or Tetrahedron, Nol. 51, p. 5813 (1995).

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wherein R⁵ represents hydroxyl group or an alkyl group; R⁶ represents chlorine atom or an alkyl group; R¹ has the same meaning as that defined for R¹ (except for hydroxyl group); and R³, m and ring A have the same meanings as those defined above.

[0027] In Step 1, the compound of the general formula (IV) can be obtained by allowing the compound represented by the general formula (III) to react with a nitrating agent such as concentrated nitric acid and furning nitric acid in the presence or absence of acetic acid, sulfuric acid or the like at a temperature ranging from 0°C to 200°C.

[0028] In Step 2, the compound of the general formula (V) can be obtained by allowing the compound of the general formula (IV) to react with an appropriate chlorinating agent, for example, phosphorus exychloride, thlory/ chloride, phosphorus pentachloride or the like, in the presence or absence of a solvent such as toluene at a temperature ranging from 0°C to 200°C.

[0029] In Step 3, the compound of the general formula (VII) can be obtained by reacting the amine represented by the general formula (VI) with the compound of the general formula (V) in a solvent such as N,N-dimethylformamide and foluene in the presence or absence of a base such as triethylamine and potassium carbonate at a temperature ranging from -10°C to the reflux temperature of a solvent.

(0030) In Step 4, the compound of the general formula (VIII) can be obtained by reducing the nitro group in the compound of the general formula (VIII) according to an appropriate reducing method, for example, catalytic reduction using a metal catalyst such as platinum, Raney nickel, and palladium/carbon; reduction using nickel chloride and so-dium borohydride; reduction using nickel chloride and nickel nickel

[0031] The reduction can be carried out in a solvent such as water, methanol, ethanol, and tetrahydrofuran, as well as a mixed solvent thereof, at a temperature ranging from 0°C to the reflux temperature of the solvent.

[0032] In Step 5, the compound of the general formula (IX) can be obtained by reacting the compound of the general formula (VIII) with a compound represented by the following general formula (XI), (XII) or (XIII):

wherein R represents a lower alkyl group; X represents a halogen atom; R1' has the same meaning as that defined for R1 (except for hydroxyl group),

in the presence or absence of a basic catalyst such as triethylamine, or an acid catalyst such as hydrochloric acid and p-tolurenesulfonic acid, in the presence or absence of a solvent such as N,N-dimethylformamide, tetrahydrofuran, actionitrile, xylene and toluren, at a temperature ranging from 0°C to 200°C.

[0033] In Step 6, as a method in place of Step 5, the compound of the general formula (IX) can be obtained by or reacting the compound of the general formula (VIII) with a compound represented by the following general formula (XIV):

wherein R¹ has the same meaning as that defined for R¹ (except for hydroxyl group), in the presence o12,3-dichloro-5,6-digyano-1,4-benzoquinone in a solvent such as acetonitrite, 1,4-dioxane and tetrahydrofuran at a temperature ranging from 070 to the refulx imperature for the solvent.

[0034] In Step 7, as a method in place of Step 5 or 6, the compound of the general formula (X) can be obtained by reacting the compound of the aforementioned general formula (VIII) with a compound represented by the following general formula (XV):

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wherein R¹ has the same meaning as that defined for R¹ (accept for hydroxyll group), in the presence or absence of an acid catalyst such as hydrochloric acid and sulfuric acid, in the presence or absence of a solvent such as NNdimethyllormamide and toluene, at a temperature ranging from 0°C to 200°C. Moreover, when R² represents hydroxyl group in the general formula (X), the compound of the general formula (IX) can be obtained by carrying out chlorination in Step 8.

[0035] The chlorination is carried out by protecting the compound of the general formula (X), if desired, at the nitrogen atom not bound to the (CP_b)_m group, that is adjacent to the saturated integen-containing heterocyclic group represented by FR, with a protecting group such as alkanoyl groups in a conventional manner, then reacting with an appropriate chlorinating agent, for example, phosphorus oxychloride, thionyt chloride, phospene, oxaly/chloride, phosphorus pentachloride or the like in the presence or absence of a selvent such as tolkene at a temperature ranging from °C to 20°C, and further deprotecting in a conventional manner, if desired, to obtain the compound of the general formula (X) wherein R8 is chlorina story.

[0036] In the second synthetic method of the compounds of the present invention, the compound of the general formula (XVI):

wherein R², R⁸, m and ring A have the same meanings as those defined above, can be obtained by allowing the compound of the general formula (VIII) to react together with triphosgene in the presence of a base such as triethylamine and potassium carbonate in a solvent such as 1,2-dichloroethane, 1,4-dioxane, tertrahydrofuran, N.N-dimethylformamide and toluene at a temperature ranging from 0°C to the reflux temperature of a solvent.

[0037] In the third synthetic method of the compounds of the present invention, the compound of the general formula (XVII):

$$\begin{array}{c} \text{CH}_3\\ \text{Z-S(O)}_a\\ \text{A}\\ \text{N}\\ \text{R}^8 \end{array} (\text{XVIII}$$

wherein Z represents an aromatic ring; the symbol "a" represents an integer of 1 or 2; and \mathbb{R}^3 , \mathbb{R}^6 , \mathbb{R} , \mathbb{R} and ring A have the same meanings as those defined above, can be obtained by carrying out suitable oxidation of the compound of the general formula (IX) which has an any group substituted with methything group as \mathbb{R}^4 , after protecting if desired, the nitrogen atom not bound to the (CH₂)_m group, that is adjacent to the saturated nitrogen-containing heterocyclic group represented by \mathbb{R}^3 , with a protecting group such as alkanoyl groups in a conventional manner, if desired.

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[0038] The oxidation can be carried out in various manners according to the desired product. More specifically, the preparation can be made, when the symbol "a" represents an integer of 1, by reacting with an oxidizing agent, for example, chromic acid, hydrogen peroxide, m-chloroperbenzoic acid, sodium periodate, potassium periodate or the like, or when the symbol "a" represents an integer of 2, with an oxidizing agent, for example, chromic acid, hydrogen peroxide, m-chloroperbenzoic acid, senitum tetrausside; unthenium tetrausside or the tiles, in a solvent such as tetrahy-drofuran, 1.4-dioxane, 1.2-dichloroethane, methanol, acctone, and water, as well as a mixed solvent thereof, at a temperature ranging from 0"C to the refulx temperature of a solvent.

[0039]. In the form synthetic method of the compounds of the present invention, the compound of the general formula (by wherein R² is hydroxyl group can be obtained by allowing a compound of the general formula (by wherein R² is chlorine atom to react with water and an appropriate acid or base in a solvent at a temperature ranging from 0°C to the reflux temperature of a solvent. Examples of the appropriate acid include, for example, organic acids such as formic acid, acet eacil, and trifluoroacetic acid, and mineral acids such as hydrochloric acid; suffund acid, and hydrobromic acid. Examples of the appropriate base include, for example, hydroxides, carbonates and hydrogenocarbonates of alkali metal such as sodium and potassium and of alkina fine-earth metal such as smapsissum and calcium and the like. Examples of the solvent include, for example, alcohols such as methanol, alhanol and n-propen, N,N-dimethylformamidia. 1.4-dioxen, tetrahydrotrum and the like. Examples of the solvent include, for example, alcohols such as methanol, alhanol and n-propen.

[0040] In the fifth synthetic method of the compounds of the present invention, the compound of the general formula (i) wherein R² is fluorine atom, bromine atom or todine atom and R² is R² can be obtained by allowing a compound which is obtained by reacting the compound of the general formula (i) wherein R² is chlorine atom and R¹ is R² or wherein R² is hydroxyl group and R² is R² with trifluoromethanesulfonic anhydride, methanesulfonyl chloride or ptoluenesulfonyl chloride to react with a metal halidle (e.g., potassium fluoride, sodium fluoride, lithium fluoride, potassium bromide, sodium bromide, potassium loidide, sodium loidide, etc.) in an aproide solvent such as dimethylsulfoxide, N, M-dimethylformamide, and acetomicifie in the presence or absence of a phase-transfer catalyst such as tetraphenylphosphonium bromide, hexadecyliributylphosphonium bromide, and 18-crown-6 at a temperature ranging from 0°C to the reflux temperature of a solvient.

[D041] In the sixth synthetic method of the compounds of the present invention, the compound of the general formula (i), wherein R² is a saturated integen-containing heterocycle group of which the nitrogen stom that is not bound to the adjacent (CH₂)_m group is deprotected, can be obtained by subjecting the compound of the general formula (i), wherein R² is a saturated nitrogen-containing heterocyclic group having a protecting group such as alkanoy; groups, alkoxycathonyl groups, benzy group and trifluoromethyl group on the nitrogen atom which is not bound to the adjacent (CH₂)_m group, to deprotection with an acid or alkali, or to catalytic reduction with a metal catalyst, according to the type of the protecting group of the nitrogen atom.

[0042] The deprotection by using an acid or alkali can be carried out with an appropriate acid or base in the presence or absence of a cation scavenger such as anisola and tibicanisole in a solvent. Examples of the solvent used include, for example, ethyl acetate, methylene chloride, 1,2-dichlorositane, 1,4-dioxane, methanol, ethanol, p-propanel, NN-dimethylformamide, tetrahydrofuran, and water, as well as a mixed solvent thereof. Examples of the acid used include, for example, hydrochloric acid, an ethyl acetate solution of hydrogen chloride, an ethanolic solution of hydrogen chloride, suffurior acid, hydrotromic acid, tilluorosacetic acid, methanosufforic acid, p-forumental colic, for example, hydroxides, carbonates and hydrogen chlorides of alkali metal such as sodium and potassium, and of alkaline-earth metal such as magnesium and calcium and the like. The reaction can be cerried out at a temperature ranging from 0°C to the reflux emperature of a solvent.

[0043] The catalytic reduction can be carried out by using an appropriate metal catalyst such as platinum, palladium/

carbon, Raney nickel, Peariman's reagent in water, an alcohol such as methanol, ethanol and n-propanol, and acetic acid, as well as a mixed solvent thereof in the presence or absence of an acid such as hydrochloric acid at a temperature ranging from room temperature to the reflux temperature of the solvent under a pressure ranging from normal pressure to 200 kg/cm².

- 5 [0044] In the seventh synthetic method of the compounds of the present invention, the compound of the general formula (1) wherein R2 is phenoxy group which may be substituted can be obtained by reacting the compound of the general formula (1) wherein R2 is chlorine atom with a phenol derivative which may be substituted in the presence of a base such as sodium hydroxide and potassium hydroxide in the presence or absence of a solvent such as N,N-dimethylofimamide and toluene at a temperature randing from °CC to 200°C.
- 10 [0045] In the eighth synthetic method of the compounds of the present invention, the compound of the general formula (f) wherein R2 is amino group can be obtained by subjecting the compound of the general formula (f) wherein R2 is phenoxy group which may be substituted, that is obtained by the seventh synthetic method, to reaction together with ammonium acetals in the presence or absence of a solvent such as N,N-dimethylformamide and toluene at a temperature ranging from 0°C to 200°C.
- (9 Q04G) In the ninth synthetic method of the compounds of the present invention, the compound of the general formula (1) wherein R² is amino group which may have one or two substituents or a cyclic amino group which may be substituted can be obtained by subjecting the compound of the general formula (1) wherein R² is otherine atom to reaction together with an amine derivative which may have one or two substituents or a cyclic amine derivative which may be substituted in the presence or absence of a base such as triethylamine, potassium carbonate and sodium hydride in the presence or absence of a solvent such as water, alcohols including methanol, ethanol and n-propanol, methylene chloride, 1,2-dichitrochane, NN-dimethylformamide, 1,4-dioxane, tetrahydrofuran and toluene at a temperature ranging from 0°C to 20°C under normal pressure or a pressured on online.
- [0047] In the tenth synthetic method of the compounds of the present invention, the compound of the general formula (i) wherein R² is amino group can be obtained by subjecting the compound of the general formula (i) wherein R² is benzylamino group, othersylamino group, othersylamino group, which is obtained in the ninth synthetic method, to catalytic reduction by using an appropriate metal catalyst, or by subjecting the compound of the general formula (i) wherein R² is prehitosybenzylamino group to deprotection using an acid.
- [0048] The catalytic reduction can be carried out with a metal catalyst such as palladium/carbon and Peariman's reagent in a solvent such as alcohols including methanol and ethanol, and water, as well as a mixed solvent thereof?

 at a temperature ranging from room temperature to the reflux temperature of a solvent in the presence or also and discount and expensive ranging from room temperature to the reflux temperature of a solvent in the presence or also and cyclohexane, and cyclohexane, and cyclohexane under a pressure ranging from normal pressure to 200 kg/cm². The deprotection using an acid can be carried out with an acid such as hydrochloric acid, sulfuric acid, triflutoracetic acid and triflutoromethanesulfonic acid in a solvent such as alcohols including methanol and ethanol, methylene children, 1.2-dichlorethane, 1.4-dioxane, tetrahydrofuran, set of the control of the co
- [0049] In the eleventh synthetic method of the compounds of the present invention, the compound of the general formula (I) wherein R³ is a saturated nitrogen-containing heterocyclic group which is substituted with oxo group can be obtained by reacting the compound of the general formula (I) wherein R³ is a saturated nitrogen-containing heterorycolic group which is substituted with ethylenedioxy group, with an acid such as hydrochloric acid, an ethyl acetate solution of hydrogen chioride, sulfure acid, hydrobromic acid, firmino acid, firminomic acid, firmino
- 5 [0050] In the twelfth synthetic method of the compounds of the present invention, the compound of the general formula (1) wherein R³ is a saturated nitrogen-containing heterocyclic group which is substituted with hydroxylmino group can a kookylmino group can be obtained by reacting the compound of the general formula (1) wherein R³ is a saturated nitrogen-containing heterocyclic group which is substituted with oxo group, that is obtained by the eleventh synthetic method, with a compound represented by the following general formula (XVIII):

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$$R^7$$
-O-NH₂ (XVIII)

wherein R² represents hydrogen atom or an alkyl group, in the presence or absence of a base such as it intellylamine, disopropylethylamine, sodium carbonate, potassium carbonate, sodium hydrogencarbonate and sodium acetate in a solvent such as alcohols including methanol, ethanol and n-prosenol, N-d-imethylformamids 1.4 doksone, tetahydrojura, and tolevene at a temporature raminis from YC start.

to the reflux temperature of a solvent.

[0051] In the thirteenth synthetic method of the compounds of the present invention, the compound of the general formula (i) wherein R² is chlorine atom can be obtained by subjecting the compound of the general formula (i) wherein R² is chlorine atom to catalyfic reduction using a metal catalyst such as platinum and palladium/carbon in the presence or absence of an acid such as hydrochloric acid and acetic acid in an alcohol solvent such as methanol and ethanol or a water-containing solvent thereof under normal pressure at a temperature ranging from room temperature to the reflux temperature of a solvent.

[0052] In the fourteenth synthetic method of the compounds of the present invention, the compound of the general formula (i), wherein P3 is a saturated nitrogen-containing heterocyclic group having an appropriate substituent on the nitrogen atom which is not bound to the adjacent (C1-½)_m group, can be obtained by reacting an appropriate reagent with the compound of the general formula (i) wherein P3 is a saturated nitrogen-containing heterocyclic group not having a protecting group on the nitrogen atom which is not bound to the adjacent (C1-½)_m group.

[0053] The reaction can be carried out in the presence or absence of a solvent such as N,N-dimethylformamide, methylene chloride, tetahydroffuran, toluene, pyridine, nitrobernzene, 1,2-dichloroethane, 1,4-dioxane, methanol, eth-anol, propanol and water, as well as a mixed solvent thereof, in the presence or absence of a base such as triethyl-amine and potassium carbonate at a temperature ranging from 0°C to 200°C.

[0054] Examples of the appropriate reagent include, for example, alkyl halides, triphenylmethyl chloride, benzyl chloride, ethyl chlorocarbonate, di-tert-butyl dicarbonate, sodium cyanate, alkyl socyanates, sodium thiocyanate, alkyl isothiocyanates, 1H-pyrazole-1-carboxamidine, methanesulfonyl chloride, p-tucrobenzenesulfonyl chloride, urethanes, alkyltrethanes, thiourethanes, alkyltrethanes, thiourethanes, alkyltrethanes, thiourethanes, alkyltrethanes, thiourethanes, alkyltrethanes, thiourethanes, alkyltrethanesulfonyl chloride, p-tucrobenzenesulfonyl chloride, urethanesulfonyl chloride, urethanesulfonyl chloride, p-tucrobenzenesulfonyl chloride, urethanesulfonyl chloride, p-tucrobenzenesulfonyl chloride, urethanesulfonyl chloride, urethanesulfonyl chloride, p-tucrobenzenesulfonyl chloride, urethanesulfonyl chloride, urethanesulfonyl chloride, urethanesulfonyl chloride, p-tucrobenzenesulfonyl chloride, urethanesulfonyl chloride,

[0055] In the fifteenth synthetic method of the compounds of the present invention, the compound of the general formula (I), wherein R³ is a saturated nitrogen-containing heterocyclic group substituted with an alkoxycarbonyl group or benzyloxycarbonyl group on the nitrogen atom which is not bound to the adjacent (CH₂)_m group, can be obtained by reacting the compound of the general formula (I) wherein R³ is a saturated nitrogen-containing heterocyclic group substituted with an alkyl group or the nitrogen atom which is not bound to the adjacent (CH₂)_m group with an alkyl chiorocarbonate or benzyl chiorocarbonate in the presence or absence of a solvent such as methylene chioride and toluene in the presence or absence of a base such as triethylamine and potassium carbonate at a temperature ranqing from 0°C to 200°C.

[0056] Some of the compounds represented by the general formulas (III) to (VIII) which are starting materials or synthetic intermediates in the preparations of the compounds of the present invention are known compounds, which are disclosed in, for example, Journal of Medicinal Chemistry, Vol. 18, p. 726 (1975; Vol. 33, p. 1860 (1990); and Vol. 40, p. 1779 (1997); International Patent Publication No. 97/20820; European Patent Publication No. 223124 (1987) and the like, and can be prepared according to the method described therein. The preparations of some novel compounds will be described in reference examples.

[0057] The medicaments which comprise as an active ingredient the novel 1H-imidazopyridine derivative represented by the aforementioned general formula (I) or (II) or a salt thereof are generally administered as oral preparations in the forms of capsules, tablets, fine granules, granules, powders, syrups, dry syrups and the like, or as parenteral preparations in the forms of injections, suppositories, eye drops, eye ointments, ear drops, nasal drops, dermal preparations. inhalations and the like. These formulations can be manufactured according to conventional methods by addition of pharmacologically and pharmaceutically acceptable additives. For example, in the oral preparations and suppositories, pharmaceutical ingredients may be used such as excipients such as lactose. D-mannitol, corn starch, and crystalline cellulose; disintegrators such as carboxymethylcellulose and carboxymethylcellulose calcium; binders such as hydroxvpropylcellulose, hydroxypropylmethylcellulose, and polyvinylpyrrolidone; jubricants such as magnesium stearate and talc; coating agents such as hydroxypropylmethylcellulose, sucrose, and titanium oxide; bases such as polyethylene glycol and hard fat and the like. In injections, or eye or ear drops and the like, pharmaceutical ingredients may be used such as solubilizers or solubilizing aids which may constitute aqueous preparations or those dissolved upon use such as distilled water for injection, physiological saline, and propylene glycol; pH modifiers such as inorganic or organic acids or bases, isotonicities such as sodium chloride, glucose, and glycerin; stabilizers and the like; and in eye ointments and dermal preparations, pharmaceutical ingredients which are suitable for ointments, creams and patches such as white vaseline, macrogols, glycerin, and cotton cloth,

[0058] A dose of the compounds of the present invention to a patient under therapeutic treatment is generally from about 0.1 to 1,000 mg in oral administration, and from about 0.01 to 500 mg in parenteral administration for an adult, which may depend on the symptoms of the patient. The aforementioned dose can be administrated once a day or several times a day as divided portions. However, it is desirable that the aforementioned dose may suitably be increased or decreased according to a purpose of a therapeutic or preventive treatment, part or type of a disease, and the age or symptoms of a patient.

Examples

[0059] The present invention will be explained by referring to Reference Examples and Working Examples. However, the scope of the present invention is not limited to these examples.

[0060] The abbreviations in the tables have the following meanings: Ph, phenyl; Bn, benzyl; Boc, tert-butoxycarbonyl; Ac, acetyl; Ms, methanesulfonyl; Ts, p-toluenesulfonyl; Me, methyl; Et, ethyl; n-Bu, n-butyl.

Reference example 1

10 Ethyl N-triphenylmethyl-4-piperidinecarboxylate

[0061] To a solution of 78.5 g of ethyl isonipecotate and 81.5 ml of triethylarmine in 750 ml of methylene chloride, 149 g of triphenylmethyl chloride divided in three portions was added portionwise a 160 portion work and the mixture was stirred for 16 hours. The reaction mixture was added with water and extracted with methylene chloride. The extract was washed successively with water and esturated brine, and dried, and then the solvent was evaporated. The resulting brown liquid was added with disporpoyl other, and the precipitated crystals were collected by filteration and washed with disporpoyl other to give 184 g of pale yellow crystals. Recrystallization from ethanol gave coloriess prisms having the melting point of from 147.5 to 148.5°C.

Elemental analysis for C ₂₇ H ₂₉ NO ₂				
Calculated %				
Found %	C, 81.19;	H, 7.22;	N, 3.44	

25 Reference example 2

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N-Triphenvlmethyl-4-piperidinemethanol

[0062] To a suspension of 10.6 g of lithium aluminium hydride in 300 ml of dried tetrahydrofuran, a solution of 112 g of ethyl N-triphenylmethyl-4-pendidine-cateopylate in 400 ml of dried tetrahydrofuran was added droywise under ienceoling, and the mixture was stirred at room temperature for 4 hours. The reaction mixture was added dropwise with a mixture of tetrahydrofuran and 10% aqueous sodium hydroxide solution under loc-cooling, An insoluble matter was filtered off and washed with interhydrofuran. The littlates were combined and concentrated to give a coloriese solid. The colorioses solid was washed with methanol to give 84.2 g of colorides crystals. Recrystallization from methanol gave coloriese solid was cystale having the methiag point of from 92 to 95.5°C.

Elemental analysis for C ₂₅ H ₂₇ NO						
	Calculated %	C. 83.99;	H, 7.61;	N, 3.92		
	Found %	C, 83.79;	H, 7.74;	N, 3.94		

[0063] In accordance with the method of Reference example 2, the compound of Reference example 3 was obtained.

Reference example 3

N-Triphenylmethyl-4-piperidineethanol

[0064]

Appearance: colorless liquid
NNR spectrum 8 (CDOL)ppm: 126(1H,brs), 1.36(2H,brs), 1.45-1.58(4H,m), 1.67(2H,d, J=12+tz), 3.05(2H brs), 3.74(2H,L)=8+tz), 7.14(3H,L)=7.5Hz), 7.24(6H,L)=7.5Hz), 7.46(6H,brs)
IR spectrum 7 (In)_am⁻¹, 2416

Mass spectrum m/z: 371(M+)

Reference example 4

(N-Triphenylmethyl-4-piperidyl)methyl methanesulfonate

9 [0065] To a solution of 84.0 g of N-tripheny/methyl-4-piper/dimensethanol and 36.2 m for triently/amine in 420 m for direle terrary/dorfuran, 18.3 m for methanesulamyl chindrie was added droywise under loe-cooling, and the mixture was stirred at room temperature for 5.5 hours. The reaction mixture was added dwith water and extracted with dethyl other. The extract was washed successively with water and saturated brine, and dried, and then the solvent was evaporated. The resulting residue was added with a mixture of isopropanol and methanol, and the precipitated crystals were collected by filtration and washed with methanol to give 90.4 g of colorloss crystals, Recrystallization from a mixture of methylene chloride and methanol gave coloriess prisms having the melting point of from 126.5 to 134°C.

Elemental analysis for C ₂₆ H ₂₉ NO ₃ S				
Calculated %	C, 71.69;	H, 6.71;	N, 3.22	
Found %	C, 71.68;	H, 6.47;	N, 3.19	

[0066] In accordance with the method of Reference example 4, the compound of Reference example 5 was obtained.

20 Reference example 5

2-(N-Triphenvlmethyl-4-piperidyl)ethyl methanesulfonate

[0067]

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Appearance: colorless crystals
Recrystallization solvent: methanol - diethyl ether

mp: 111.5-114°C

Elemental analysis for C ₂₇ H ₃₁ NO ₃ S					
Calculated %					
Found %	C, 72.03;	H, 7.12;	N, 3.14		

35 Reference example 6

4-Azidomethyl-N-triphenylmethylpiperidine

[0068] A suspension of 60.0 g of (N-triphenylmethyl-4-piperidylymethyl methanesulfonate and 17.9 g of sodium azide in 300 ml of dried NI-dimethyl-formamide was stirred at 70°C for 17 hours. After the reaction, an insoluble matter was filtered off and the filtrate was concentrated. The resulting residue was added with water and extracted with ethyl acetate. The extract was washed successively with water and saturated brine, and dried, and then the solvent was evaporated. The resulting solid was washed successively with ethanel and n-havener to give 42.6 g of coloriess crystate. Recrystallization from a mixture of methanol and diethyl ether gave coloriess crystals having the melting point of from 103.5 to 10.5 °C.

Elemental analysis for C ₂₅ H ₂₆ N ₄					
Calculated % Found %	C, 78.50; C, 78.45;				

Reference example 7

tert-Butyl 2-(2-azidoethyl)-1-piperidinecarboxylate

[0069] To a solution of 46.7 g of tert-butyl 2-(2-hydroxyethyl)-1-piperidine-carboxylate and 31.3 ml of triethylamine in 300 ml of dried tetrahydrofuran, 15.8 ml of methanesulfonyl chloride was added dropwise under ice-cooling, and the mixture was stirred at room temperature for 2 hours. The reaction mixture was added with water and extracted with

diethyl ether. The extract was washed successively with water and saturated brine, and dried, and then the solvent was evaporated. The resulting solid was washed with n-heptane to give 6-4, 9 of colorless crystals. And then, 22-9 g of sodium azide and 220 mil of N./- Meinethylformanide were added to the resulting crystals, and the mixture was stirred at 70°C for 4 hours. After the reaction, an insoluble matter was filtered off and the filtrate was concentrated. The resulting residue was added with water and extracted with ethyl accitate. The extract was washed successively with water and saturated brine, and dried, and then the solvent twas evaporated to give 43.2 g of a yellow liquid.

NMR spectrum & (DMSO-d_s)ppm: 1.20-1.32(1H,m),1.40(9H,s),1.48-1.58(5H,m),1.60-1.68(1H,m),1.88-1.96(1H, m),2.71-2.78(1H,m),3.28(2H;I,J-6.5Hz),3.80-3.88(1H,m),4.19-4.25(1H,m) IR spectrum v (ilia.)tm⁻¹: 2104.1692

Reference example 8

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4-Oxo-1-piperidineacetonitrile

[0070] A suspension of 25.0 g of 4-piperidinone monohydrochloride monohydrate, 11.5 ml of chloroacetonitrile and 57.0 ml of dilisopropylethylamine in 250 ml of tetrahydrotran was refluxed for 10 hours. After the reaction, an insoluble matter was filtered off. The filtrate was added with saturated aqueous sodium hydrogenoarbonate solution and extracted with a mixture of eithyl acetate and methanol (10:1). The extract was dried, and the solvent was evaporated to give brown crystals. The crystals were washed with a mixture of eithyl acetate and n-heptane to give 15.7 g of pale brown crystals.

NMR spectrum δ (CDCl3)ppm: 2.53(4H,t,J=6Hz),2.91(4H,t,J=6Hz),3.66(2H,s) IR spectrum v (KBr)cm^1: 2232,1714

Mass spectrum m/z: 138(M+)

[0071] In accordance with the method of Reference example 8, the compound of Reference example 9 was obtained.

Reference example 9

4-(tert-Butoxycarbonylamino)-1-piperidineacetonitrile

[0072]

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Appearance: colorless needles Recrystallization solvent: methanol mp: 147-148°C

Elemental analysis for C ₁₂ H ₂₁ N ₃ O ₂				
Calculated %				
Found %	C, 60.08;	H, 8.63;	N, 17.55	

Reference example 10

N-Triphenylmethyl-4-piperidineacetonitrile

[0073] A suspension of 90.4 g of (N-triphenyhrethyl-4-piperidy)/methyl methanesulfonato, 3.50 g of potassium iodide and 20.3 g of sodium cyanide in 400 m tol dried dimethylsulfloxide was stirred at 90°C for 5 hours. The reaction mixture was action with water and extracted with eithyl acetate. The extract was washed successively with water and saturated brine, and dried, and the solvent was evaporated to give a yellow liquid. The liquid was added with methanol, and the precipitated crystals were collected by filtration and washed with methanol to give 7.0 g of coloriess crystals. Recrystallization from a mixture of methylene chloride and methanol give coloriess crystals having the melting point of from 38 to 139°C.

Elemental analysis for C ₂₆ H ₂₆ N ₂					
Calculated %	C, 85.21;	H, 7.15;	N, 7.64		
Found %	C, 85.35;	H, 7.26;	N, 7.62		

[0074] In accordance with the method of Reference example 10, the compounds of Reference examples 11 through 13 were obtained.

Reference		Physical properties
example		(Recrystallization solvent)
		colorless crystals (MeOH-Et ₂ O)
		mp,158.5−160.5°C
11	Ph ₃ CN	Elemental analysis for C ₂₇ H ₂₈ N ₂
	CN	Caled. %: C, 85.22; H, 7.42; N, 7.36
		Found %: C, 85.21; H, 7.52; N, 7.34
		colorless prisms (iso-Pr ₂ O-n-Heptane)
	Bock	mp,48-49°C
12		Elemental analysis for C ₁₂ H ₂₀ N ₂ O ₂
		Calcd. %: C, 64.26; H, 8.99; N, 12.49
		Found %: C, 64.01; H, 9.24; N, 12.35
		colorless crystals (iso-Pr ₂ 0)
		mp,89−90°C
13	Boch. CN	Elemental analysis for C ₁₁ H ₁₈ N ₂ O ₃
	BOUR	Calcd. %: C, 58.39; H, 8.02; N, 12.38
		Found %: C, 58.31; H, 8.01; N, 12.37

Reference example 14

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N-Triphenylmethyl-4-piperidineacetic acid

[0075] A suspension of 21.2 g of N-triphenylmethyl-4-piperidineacetonitrile, 127 ml of 10% aqueous sodium hydroxide solution and 312 ml of ethanol was refluxed for 74 hours. The reaction mixture was neutralized with 10 % hydrochloric acid under ice-cooling, and then adjusted to pH 4-5 with 10% aqueous citric acid solution. The precipitated crystals were collected by filtration, and washed successively with water and methanol to give 23.6 g of coloriess crystals. Recrystallization from a mixture of methanol and ethyl acetate gave coloriess needles having the melting point of from 197 to 290°C (decomposition).

Elemental analysis for C ₂₆ H ₂₇ NO ₂				
Calculated % Found %	C, 81.01;	H, 7.06;	N, 3.63	
Found %	C, 80.85;	H, 7.17;	N, 3.70	

Reference example 15

Ethyl N-triphenylmethyl-4-piperidineacetate

9 [0076] A suspension of 23.8 g of N-triphenymethyl-4-piperidineacotic acid, 16.9 g of potassium cerbonate and 5.0 mixture was added with water and 6thyl bronide in 230 ml of dried N.N-dimethyllomanide was stirred at 90°C for 5 hours. After cooling, the reaction mixture was added with water and 6thyl acetate, and the precipitated crystals were collected by filtration and washed with water to give 20.6 g of colorless crystals. Recrystallization from a mixture of methanol and tetrahydrofuran gave colorless crystals having the metiting point of from 185 to 168°C.

Elemental analysis for C ₂₈ H ₃₁ NO ₂				
Calculated %				
Found %	C, 81.08;	H, 7.69;	N, 3.43	

Reference example 16

4.4-Ethylenedioxy-1-piperidineacetonitrile

Q077] A solution of 1.00 g of 4-oxo-1-pleridineactiontrile, 22.6 g of ethylene glycol and 0.62 g of anhydrous p-toluenesulfonic acid in 100 ml of toluene was refluxed for 6 hours with Dean-stark dehydrating apparatus. After cooling, the reaction mixture was added with saturated aqueous sodium hydrogencarbonate solution and extracted with ethyl acetate. The extract was dried, and the solvent was evaporated to give a pale brown liquid. The resulting liquid was purified by alumina column chromatography using ethyl acetate - n-heptane (1:3) as an eluting solvent to give 12.8 g s of a colorless flouid.

NMR spectrum 6 (CDCl₃)ppm: 1.78(4H,t,J=6Hz),2.69(4H,t,J=6Hz),3.52(2H,s),3.96(4 H,s) IR spectrum v (i(1,9m⁻¹: 2230.1094 Mass spectrum m/z: 182(M⁻¹)

Reference example 17

4-Aminomethyl-N-triphenylmethylpiperidine

5 [0078] To a suspension of 4.70 g of lithium aluminium hydride in 250 ml of dried tetrahydrofuran, a solution of 47.7 g of 4-azidomethyl-h-triphenyimethyloperidine in 250 ml of dried tetrahydrofuran was added dropwise under lex-cooling, and the mixture was atded dropwise with a mixture of tetrahydrofuran and 10% aqueous sodium hydroxide solution under ice-cooling. An insolutio matter in the mixture was affiltered off, and washed with letrahydrofuran. The filtrate and the washings were combined and concentrated to give 48.1 of a colorless flouid.

NMR spectrum 6 (CDCl₉)ppm: 1.14(1H,brs), 1.36(2H,brs), 1.48(2H,qd,J=5,2.5Hz), 1.68 (2H,d,J=11.5Hz), 2.59(2H,d,J=6Hz), 3.10(2H,brs), 7.14(3H,LJ=7,5Hz), 7.25(6H,LJ=7,5Hz), 7.47(6H,brs) (IR,port); 0.656,3028

High resolution mass spectrum: Analysis for C25H28N2

Calculated m/z: 356.2252 Found m/z: 356.2250

50 Reference example 18

4-(2-Aminoethyl)-N-triphenylmethylpiperidine

[0079] To a suspension of 21.7 g of lithium aluminium hydride in 300 m lof dried letrahydrofuran, a solution of 28.1 g of concentrated sulfuric acid in 100 m lof dried tetrahydrofuran was added dropwise under ice-cooling, and the mixture was stirred for 30 minutes. And then, a solution of 70 0 g of N-triphenylmethyl-4-piperidimeacetonitrie in 300 m lof dried tetrahydrofuran was added dropwise to the mixture was role of the mixture was retired at room temperature for 8 hours. The reaction mixture was added dropowise with a mixture of tetrahydrouran and 10% auguous sodium.

hydroxide solution under ice-coeling. An insoluble matter in the mixture was filtered off, and the filtrate was concentrated. The resulting residue was added with water and extracted with ethyl acetate. The extract was washed with saturated brine, and dried, and the solvent was evaporated to give 71.4 g of a colorless liquid.

NMR spectrum δ (CDC_b)ppm: 1.18(1H,brs),1.35(2H,brs),1.40(2H,q,J=7.5Hz),1.48(2 H,qd,J=11.5,3Hz),1.63(2H,d,J=11.5Hz),2.67(2H,I,J=7.5Hz),3.05(2H,brs),7.14(3H,I,J=7.5Hz),7.24(6H,I,J=7.5Hz),7.47(6H,brs)
IR spectrum v (iii.,2mr*,3060,3032

High resolution mass spectrum: Analysis for C26H30N2

10 Calculated m/z; 370.2409 Found m/z: 370.2400

[0080] In accordance with the method of Reference example 18, the compound of Reference example 19 was obtained.

Reference example 19

4-(3-Aminopropyl)-N-triphenylmethylpiperidine

20 [0081]

30

Appearance: colorless liquid

NMR spectrum δ (DMSO-d₀)ppm: 0.95+1.05(1H,m),1.19+1.35(6H,m),1.41(2H,q,J=11.5Hz), 1.62(2H,d,J=11.5Hz), 2.47(2H,t,J=6.5Hz),2.93(2H,d,J=11.5Hz),7.15(3H,t,J=7.5Hz),7.28(6H,t,J=7.5Hz),7.38(6H,d,J=7.5Hz),7.38(6H,d,J=7.5Hz),

Reference example 20

tert-Butyl 2-(2-aminoethyl)-1-piperidinecarboxylate

[0082] A suspension of 43.0 g of tert-buly! 2-(2-azidoethyl):1-piperidinecarboxylate and 2.15 g of 5% paliadium on carbon in 215 mi of methanol was catalytically hydrogenated at room temperature for 9 hours. After the reaction, the catalyst was filtered off, and the filtrate was concentrated to give 97.2 g of a coloriess liquid. MMB spectrum 5 (DMSO-d₂)porm: 1.20-1.30(1H,m), 1.36(9H,s), 1.45-1.56(4H,m),1.72-1.82(1H,m),2.34-2.47(2H,m),2.65-2.76(1H,m),3.18(2H,t, _36H,3),3.18(2H,t, _3

IR spectrum v (liq.)cm⁻¹: 2976,2936,1692

Reference example 21

40 1-(2-Aminoethyl)-4.4-ethylenedioxypiperidine

[0083] A suspension of 12.7 g of 4.4-ethylenedioxy1-piperidineacetonitrile, 1.3 ml of Raney nickel and 113 ml of 2% methanolic solution of ammonia was catalytically hydrogenated at room temperature under 50 atm for 20 hours. After the reaction, the catalyst was filtered off, and the filtrate was concentrated. The resulting pale green liquid was puffied by alumina column chromatography [eluting solvent: ethyl acetate →ethyl acetate - methanol (10:1)] to give 10.1 g of a coloriess isluid.

NMR spectrum δ (DMSO-d₆)ppm: 1.58(4H,t,J=6Hz),2.37(2H,t,J=6.5Hz),2.42(4H,t,J=6Hz),2.57(2H,t,J=6.5Hz),3.84 (4H.s)

IR spectrum v (liq.)cm⁻¹: 2956,2884,1094

[0084] In accordance with the method of Reference example 21, the compounds of Reference examples 22 through 25 were obtained.

	Reference example		Physical properties
10	22	Bock NH ₂	colorless liquid NMR spectrum & (DMSO-d ₄)ppm:1.02-1.12(1H,m),1 .16-1.50(14H,m),1.53-1.60(1H,m),1.70-1.77(1H,m),2. 58(2H,t,J=7.5Hz),2.75-2.83(1H,m),3.65-3.78(2H,m) IR spectrum \(\nu\) (liq.) cm ⁻¹ .2980,2936,1692
15	23	Bock NH ₂	bluish green liquid NMR spectrum δ (DMSO-d _a)ppm:1.40(9H,s),1.55-2. 00(2H,m),2.50-2.65(1H,m),2.75-2.90(1H,m),2.90-3.5 0(4H,m),3.60-3.90(3H,m) IR spectrum ν (liq.) cm ⁻¹ :1700
25	24	BocHN NH2	dark green liquid NMR spectrum δ (CDCL)ppm:1.15(2H,brs),1.45(9H, s),1.85-2.00(2H,m),2.00-2.20(2H,m),2.30-2.50(2H,m),2.60-2.95(4H,m),3.40-3.60(2H,m),4.46(1H,brs) IR spectrum ν (liq.) om ⁻¹ 3332,1692
35	25	NH ₂	colorless liquid NMR spectrum δ (DMSO-d _b)ppm:1.39(9H,s),1.58-1. 65(1H,m),1.68-1.90(5H,m),2.47(2H,t,1=7.5H2),3.13-3 22(2H,m),3.68-3.76(1H,m) R spectrum ν (liq.) cm ⁻¹ -2972,2876,1896 Specific rotation [α] ₅ ²⁰ : -54.3° (c=0.1, DMSO)

Reference example 26

40

5,7-Dichloro-6-nitrothieno[3,2-b]pyridine

[0085] A mixture of 24.8 g of 4,5-dihydro-7-hydroxy-6-nitrothieno(3,2-b)pyridine-5-one and 87 ml of phosphorus oxychloride was stirred at 60°C for 24 hours. The reaction solution was concentrated and the residue was dissolved in a mixture of methylene chloride and methanol (10:1), and then the solution was poured into water. An insoluble matter was filtered off, and the organic solvent layer was separated. Furthermore, the aqueous layer was extracted with a mixture of methylene chloride and methanol (10:1). The combined organic solvent layer was dried, and the solvent was evaporated to give brown crystals. The resulting brown crystals were purified by silica gel column chromatography using ethyl acetate --n-hexane (1:3) as an eluting solvent to give 10.6 g of pale brown crystals. Recrystallization from n-hexane aver pale brown crystals having the melting point of from 96 to 97°C.

NMR spectrum δ (CDCl₃)ppm: 7.61(1H,d,J=5.5Hz).8.07(1H,d,J=5.5Hz) IR spectrum ν (KBr)cm⁻¹: 1540,1368 Mass spectrum m/z : 248.250,252(M+,9:6:1)

[0086] In accordance with the method of Reference example 26, the compounds of Reference examples 27 through

5	Reference		Physical properties
	example		(Recrystallization solvent)
10	27	CI NO2	pale brown crystals NMR spectrum o (CDCI ₁)ppm:7.87(1H,dd,J=9,2. 5Hz),8.06(1H,d,J=9Hz),8.24(1H,d,J=2.5Hz)
15	28	Me CI NO2	brown crystals NMR spectrum ở (DMSO-d _e)ppm:2.82(3H,s),7.7 8(1H,dd,J=9,2Hz),7.96(1H,d,J=2Hz),8.05(1H,d,J=9Hz)
20	29	MeO NO2	pale brown crystals NMR spectrum & (CDCl ₂)ppm:4.01(3H,s),7.42(1H ,d,J=2.5Hz),7.55(1H,dd,J=9,2.5Hz),7.99(1H,d,J=9 Hz)
25 30	30	CI NO ₂	yellow crystals (isc-PrOH) mp.182-183°C Elemental analysis for C ₆ H ₅ Cl ₁ N ₅ O ₂ Calod. N: C, 38.37; H, 1.24; N, 17.22 Found N: C, 38.37; H, 1.02; N, 17.25
35	31	CI NO2	pale brown plates (n-Hexans) mp.84-45.°C Elemental analysis for C ₉ H ₂ (i ₁ N ₁ O ₂ calcd. S. O., 43.75; H, 3.26; N, 11.34 Found S. C., 43.77; H, 3.32; N, 11.44
45	32	CI NO2	pale yellow plates (n-Hexane) mp.94.5-95.5°C Elemental analysis for C ₀ H ₀ Ci ₂ N ₂ O ₂ Coled. %: C, 41.22; H, 2.56; N, 12.02 Found %: C, 41.12; H, 2.64; N, 12.01

Reference example 33

50

2-Chloro-3-nitro-4-[2-(N-triphenylmethyl-4-piperidyl)ethylamino]quinoline

[0087] To a solution of 22.6 g of 2.4-dichlore-3-nitroquinoline and 13.0 m lof triathylamine in 60 ml of N.N-dimethylformamide, a solution of 23.0 g of 4-(2-aminoethyl)-N-triphenylmethylpiperidine in 40 ml of N.N-dimethylformamide was added dropwise with stirring under ice-cooling. The mixture was stirred at room temperature for 1 hour. The reaction mixture was added with ethyl acetate and water. The precipitated crystals were collected by filtration, and washed successively with ethyl acetate and diethyl either to give 26.9 g of y vellow crystals. Recrystallization from a mixture of N-dimethylformamide and ethyl acetate gave yellow crystals having the melting point of from 22.5 to 231°C (decomposition).

Elemental analysis for C ₃₅ H ₃₃ ClN ₄ O ₂							
Calculated %							
Found %	C, 72.64;	H, 5.80;	N, 9.82				

[0088] In accordance with the method of Reference example 33, the compounds of Reference examples 34 through 60 were obtained.

Reference				Physical properties
example	В	R ³	m	(Recrystallization solvent)
				yellow crystals(CH ₂ Cl ₂ -iso-Pr ₂ O)
				mp,196.5-199.5°C (decomposition)
34	Ci	Ph ₃ CN	2	Elemental analysis for C ₃₅ H ₃₂ Cl ₂ N ₄ O ₂
			1	Calcd.%: C, 68.74; H, 5.27; N, 9.16
				Found %:C, 68.47; H, 5.31; N, 9.18
			T-	yellow crystals(MeOH-THF)
				mp,214.5-225°C (decomposition)
35	н	Ph ₃ CN	1	Elemental analysis for C ₂₄ H ₂₁ ClN ₄ O ₂
		<u> </u>		Calcd.%: C, 72.52; H, 5.55; N, 9.95
		i i	İ	Found %:C, 72.54; H, 5.62; N, 9.82
			3	yellow crystals(MeOH-iso-Pr ₂ O)
				mp,176.5-183°C (decomposition)
36	н	Ph ₃ CN		Elemental analysis for C ₃₈ H ₃₅ ClN ₄ O ₂
		~ \		Caled.%: C, 73.14; H, 5.97; N, 9.48
				Found %: C, 73.33; H, 6.04; N, 9.36
				yellow crystals(McOH)
			2	mp,128.5−129.5°C
37	н	BnN		Elemental analysis for C ₂₃ H ₂₅ CiN ₄ O ₂
				Calcd.%: C, 65.01; H, 5.93; N, 13.19
				Found %: C, 64.96; H, 6.03; N, 13.27
38 H				yellow crystals(AcOEt)
		2		mp,199−202°C (decomposition)
	н	Bock	0	Elemental analysis for C ₁₉ H ₂₃ ClN ₄ O ₄
		~ \		Calcd.%: C, 56.09; H, 5.70; N, 13.77
		}		Found%: C, 56.04; H, 5.69; N, 13.77

Reference	В	w	Physical properties
example	6	. "	(Recrystallization solvent)
			yellow crystals(MeOH)
			mp,189.5−190.5°C
39	CI	СН	Elemental analysis for C ₂₁ H ₂₆ Cl ₂ N ₄ O ₄
			Calcd.%: C, 53.74; H, 5.58; N, 11.94
			Found%: C, 53.61; H, 5.55; N, 11.67
			yellowish orange crystals (MeOH)
	Me	СН	mp,185-186℃
40			Elemental analysis for C ₂₂ H ₂₉ ClN ₄ O ₄
			Galcd.%: C, 58.86; H, 6.51; N, 12.48
			Found%: C, 58.72; H, 6.60; N, 12.39
	MeO	СН	yellowish orange crystals (MeOH)
			mp,183.5−184.5°C
41			Elemental analysis for C ₂₂ H ₂₉ ClN ₄ O ₅
			Calcd.%: C, 56.83; H, 6.29; N, 12.05
			Found%: C, 56.90; H, 6.34; N, 12.05
			yellow crystals(AcOEt-Et ₂ O)
		N	mp,157.5-161°C
42	н		Elemental analysis for C ₂₀ H ₂₆ ClN ₅ O ₄
			Calcd.%: C, 55.11; H, 6.01; N, 16.07
			Found%: C, 55.18; H, 6.10; N, 15.86

	Reference	R²	R³	Physical properties
5	example			(Recrystallization solvent)
-				yellow crystals(AcOEt-iso-Pr ₂ O)
				mp,133-134°C
	43	CI	Bock	Elemental analysis for C ₂₁ H ₂₇ CIN ₄ O ₄
10			_ ~ `	Calcd.%: C, 57.99; H, 6.26; N, 12.88
				Found%: C, 57.99; H, 6.34; N, 12.85
				yellow crystals(EtOH)
15			^	mp,138−138.5°C
	44	Me	BocN	Elemental analysis for C ₂₂ H ₃₀ N ₄ O ₄
				Caled,%: C, 63.75; H, 7.30; N, 13.52
20				Found%: C, 63.70; H, 7.49; N, 13.44
				yellow needles (AcOEt-n-Heptane)
				mp,148.5-149°C
25	45	CI		Elemental analysis for C ₂₁ H ₂₇ ClN ₄ O ₄
			Boc	Calcd.%: C, 57.99; H, 6.26; N, 12.88
				Found%: C, 58.04; H, 6.27; N, 12.87
30				yellow crystals(iso-Pr ₂ O)
			^	mp,121-122.5℃
	46	Cí	Bock	Elemental analysis for C ₂₁ H ₂₇ ClN ₄ O ₄
35			Bock	Calcd.%: C, 57.99; H, 6.26; N, 12.88
				Found%: C, 58.04; H, 6.32; N, 12.82
				yellow prisms (MeOH-iso-Pr ₂ O)
40			^	mp,155-157°C
	47	CI	BocN	Elemental analysis for C ₂₀ H ₂₅ ClN ₅ O ₄
				Calcd.%: C, 55.11; H, 6.01; N, 16.07
45				Found%: C, 54.92; H, 5.89; N, 16.00

R³ NH NC

Reference	R ²	R3	Physical properties
example			(Recrystallization solvent)
			yellow crystals (MeOH)
		_	mp,176.5-177.5°C
48	Cı	Boch.	Elemental analysis for C ₂₀ H ₂₅ ClN ₄ O ₅
		Bock	Calcd.%: C, 54.98; H, 5.77; N, 12.82
			Found%: C, 54.85; H, 5.76; N, 12.86
			yellow needles (AcOEt-iso-Pr ₂ O)
		BocHN.	mp,150−150.5°C
49	CI	Y)	Elemental analysis for C ₂₁ H ₂₈ ClN ₅ O ₄
		~~~	Calcd.%: C, 56.06; H, 6.27; N, 15.57
			Found%: C, 55.92; H, 6.19; N, 15.59
			yellow crystals (AcOEt)
		BocHN.	mp,151−151.5°C
50	Me		Elemental analysis for C ₂₂ H ₃₁ N ₅ O ₄
		\ <u>\</u> ''\	Calcd.%: C, 61.52; H, 7.27; N, 16.31
			Found%: C, 61.33; H, 7.14; N, 16.29
			yellow fine needles (AcOEt-iso-Pr ₂ O)
		_	mp,119.5−123°C
51	CI		Elemental analysis for C ₁₈ H ₂₁ CIN ₄ O ₄
91	U	07	1/4H₂O
		V	Calcd.%: C, 54.41; H, 5.45; N, 14.10
			Found%: C, 54.60; H, 5.45; N, 14.19

R³—(CH₂)_m NH

Reference	R ³	_ m	Physical properties
example			(Recrystallization solvent)
			yellow prisms (AcOEt-n-Heptane)
	HO、 ^		mp,121-123°C
52	Y ]	2	Elemental analysis for C ₁₆ H ₁₉ ClN ₄ O ₃
	~~~		Calcd.%: C, 54.78; H, 5.46; N, 15.97
			Found%: C, 54.70; H, 5.51; N, 15.93
			yellow crystals (MeOH)
	-^		mp,123-124℃
53	, N.	2	Elemental analysis for C ₁₅ H ₁₇ ClN ₄ O ₃
	~··<		Calcd.%: C, 53.50; H, 5.09; N, 16.64
			Found%: C, 53.44; H, 4.94; N, 16.60
			yellowish brown crystals (MeOH)
		3	mp,163-164°C
54			Elemental analysis for C ₁₅ H ₁₉ ClN ₄ O ₃
			Calcd.%: C, 54.78; H, 5.46; N, 15.97
			Found%: C, 54.79; H, 5.36; N, 15.95
			yellowish brown crystals (MeOH)
- 1	\wedge		mp,145−146°C
55	\ N.	2	Elemental analysis for C ₁₈ H ₁₉ ClN ₄ O ₂
ļ	~ (Calcd.%: C, 57.40; H, 5.72; N, 16.73
			Found%: C, 57.23; H, 5.75; N, 16.74
			yellow crystals (iso-Pr ₂ 0)
			mp,102.5-103°C
56	\n\	2	Elemental analysis for C ₁₅ H ₁₇ CIN ₄ O ₂
			Calcd.%: C, 56.16; H, 5.34; N, 17.47
			Found%: C, 56.14; H, 5.37; N, 17.41

	Reference		D
			Physical properties
5	example		(Recrystallization solvent)
			yellow prisms (iso-Pr ₂ O-n-Heptane)
		_	mp,96−98°C
		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Elemental analysis for C ₂₀ H ₂₅ CiN ₄ O ₄
10	57	Boc NO ₂	Calcd.%: C, 57.07; H, 5.99; N, 13.31
			Found%: C, 57.04; H, 5.92; N, 13.26
		₩ CI	Specific rotation
15			[\alpha] _D ²⁰ : -97.3° (c=0.1, DMSO)
			pale yellow crystals (MeOH)
		BocN	mp,135-135.5℃
20	58	NH NO2	Elemental analysis for C ₂₁ H ₃₁ ClN ₄ O ₄
			Calcd.%: C, 57.46; H, 7.12; N, 12.76
		V N° CI	Found%: C, 57.33; H, 7.15; N, 12.74
25			red liquid
			NMR spectrumδ (DMSO-d ₆)ppm:0.98(2H,q,J
		BocN	=12.5Hz),1.20-1.30(1H,m),1.41(9H,s),1.59(2H,
30		V√V _{NH}	d,J=12.5Hz),2.04(2H,quin,J=8Hz),2.60-2.72(4
	59	NO ₂	H.m),2.79(2H,t,J=8Hz),2.93(2H,t,J=8Hz),3.21(2
		√ N d₁	H,q,J=6.5Hz),3.89(2H,d,J=12.5Hz),6.52(1H,t,J
35			=6.5Hz)
			IR spectrum \$\nu(\text{liq.}) cm^-1:1688,1526,1366
			orange crystals (iso-PrOH)
40		Bocn	mp,148.5−150°C
	60	V NH NO₂	Elemental analysis for C ₁₈ H ₂₅ ClN ₄ O ₄ S
			Calcd.%: C, 51.75; H, 5.71; N, 12.71
45		N CI	Found%: C, 51.64; H, 5.80; N, 12.69
40			

Reference example 61

50 3-Amino-2-chloro-4-[2-(N-triphenylmethyl-4-piperidyl)ethylamino]quinoline

[0089] To a solution of 6.56g of nickel chloride hexahytate and 22.3 ml of methanol in 100 ml of tetrahytrofuran, 2.09 g of sodium brorbydride was added portionwise under lee-cooling, and then a suspension of 31.9 g of 2-chloro-3-nitro-4-(2-(N-triphenylmethyl-4-piperidyl)ethylaminolquinoline in 300 ml of tetrahydrofuran was added to the mixture. Successively, 8.35 g of sodium borohydride divided in four portions was added portionwise, and the mixture was silred at room temperature for 1 nour. The reaction mixture was added with 50 ml of water and an insoluble matter was filtered off, and then the extract was concentrated. The residue was added with water and oxtracted with 6thyl acotate. The extract was was destroad social and dried, and then the solvent was evaporated. The

resulting pale green liquid was solidfied with a mixture of eithyl acetate and disopropyl ether, and the solid was washed successively with isopropanol and disopropyl ether to give 20.1 g of pale green crystals. Recrystallization from isopropanol gave pale green crystals having the melting point of from 116 to 121°C.

Elemental analysis for C ₃₅ H ₃₅ CIN ₄						
Calculated % C, 76.83; H, 6.45; N, 10.24						
Found %	C, 76.74;	H, 6.54;	N, 10.17			

10 [0090] In accordance with the method of Reference example 61, the compounds of Reference examples 62 through 88 were obtained.

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E R R R R R R R R R		Reference	Γ			Physical properties
colorless crystals (EtOH) mp.197-198.5°C 2 Elemental analysis for C ₃₂ H ₃ ,Cl ₂ N ₄ Calcd.*C. 72.28; H. 5.89; N. 9.53 Foundk: C, 72.45; H. 8.17; N. 9.34 brown liquid NMR spectrum & (OMSO-d ₄)ppm:1.20-1.45(3H,m.), .49(2H,d,j=11.5Hz),1.72(2H,d,j=11.5Hz),3.18(2H,t,j=7Hz),7.14(3H,t,j=7.5Hz),7.35-7.45(8H,m),7.68(1H,d,j=8Hz) 11 = 7Hz),7.99(1H,d,j=8Hz) IR spectrum \(\nu\) (liq.) om 1:3358,3058 colorless crystals (iso-Pr ₂ O) mp.149-158°C 3 Elemental analysis for C ₃₂ H ₃₇ ClN ₄ Calcd.*C. 7.705; H. 6.85; N. 9.98 Founds: C, 76.93; H. 6.81; N. 9.97 brown liquid NMR spectrum & (ODCl ₃)ppm:1.20-1.50(3H,m),1.80(2H,d,j=11.Hz),3.49(2H,d,j=11.Hz),2.7(2H,d,j=7.5Hz),3.49(2H,a),3.7 9 (HH,t,j=7.5Hz),1.66(2H,d,j=11.Hz),1.94(2H,t,j=11.Hz),2.8(2H,d,j=11.Hz),3.7(2H,d,d,j=3.15.Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,d,j=8,1.5Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.74(1H,dd,j=3.1.5Hz),7.99(1H,dd,j=8,1.5Hz),7.99(1H,dd,j		example	В	R³	m	
NMR spectrum δ (DMSO-d _a)spm:120-1.45(3H,m),1 A9(2H,q,J=11.5Hz),1.72(2H,d,J=11.5Hz),3.18(2H,t,J A9(2H,q,J=11.5Hz),1.72(2H,d,J=11.5Hz),3.18(2H,t,J 1 = 7Hz),4.89(2H,t,J=7.5Hz),7.35-7.45(8H,m),7.68(1H,d,J=8 Hz),7.99(1H,d,J=8Hz) IR spectrum ν (liq.) cm ⁻¹ :3358,3058 colorless crystals (iso-Pr ₂ O) mp,149-158°C 3 Elemental analysis for O ₂₄ H ₂₁ ClN ₄ Calcd.%: O, 77.05; H, 6.85; N, 9.98 Founds: C, 76.93; H, 6.81; N, 9.97 brown liquid NMR spectrum δ (ODCl ₃)spm:1.20-1.50(3H,m),1.80(2H,d,J=11Hz),2.34(2H,t,J=11Hz),2.88(2H,d,J=11Hz),3.27(2H,q,J=7.5Hz),3.49(2H,s),3.7 40 BnN 85 H BnN 10 Sectrum δ (DDCl ₃)spm:1.20-1.50(3H,m),1.80(1H,t,J=8,1.5Hz),7.74(1H,td,J=8,1.5Hz),7.99(1H,td,J=8,1.5Hz),7.74(1H,td,J=8,1.5Hz),7.99(1H,td,J=8,1.5Hz),7.74(1H,td,J=8,1.5Hz),7.99(1H,td,J=8,1.5Hz),7.74(1H,td,J=8,1.5Hz),7.99(1H,td,J=8,1.5Hz),7.74(1H,td,J=8,1.5Hz),7.99(1H,td,J=8,1.5Hz),7.74(1H,td,J=8,1.5Hz),7.99(1H,td,J=8,1.5Hz),7.74(1H,td,		62	CI	Ph ₃ CN	2	mp,197–198.5°C Elemental analysis for C ₃₃ H ₂₄ Cl ₂ N ₄ Calcd.½: C, 72.28; H, 5.89; N, 9.63
mp, 149-158°C 3 Elemental analysis for C ₂₈ H ₃₇ ClN ₄ Calcd.½: C, 77.05; H, 8.85; N, 9.98 Found½: C, 76.93; H, 6.81; N, 9.97 brown liquid NMR spectrum δ (CDCl ₃)ppm:1.20-1.50(3H,m),1.80(2H,d,J=1Hz),1.94(2H,J,J=1Hz),288(2H,d,J=1Hz),3.97(2H,d,J=7.5Hz),3.49(2H,J),3.97 40 41 42 43 44 45 46 47 48 48 49 49 49 40 40 40 40 41 41 42 41 43 41 43 41 43 41 44 45 41 46 46 47 48 48 49 49 40 40 40 40 41 41 42 41 43 41 43 41 43 41 43 41 43 41 43 41 43 43 43 44 44 45 46 46 47 48 48 48 49 49 40 40 40 40 40 40 40 40 40 40 40 40 40		63	Н	Ph ₃ cN	1	NMR spectrum & (DMSO-d _a)ppm:1.20-1.45(3H,m,); A9(2H,q,J=11.5Hz),1.72(2H,d,J=11.5Hz),3.18(2H,±,J=7Hz),4.89(2H,s),5.09(1H,t,J=7Hz),7.14(3H,t,J=7.5Hz),7.27(6H,t,J=7.5Hz),7.35-7.45(6H,m),7.66(1H,d,J=8Hz),7.99(1H,d,J=8Hz)
NMR spectrum δ (ODCI ₃)ppm:1.20-1.50(3H,m),1.80(2H,q,J=7.5Hz),1.86(2H,d,J=11Hz),1.94(2H,t,J=11Hz), 2.88(2H,d,J=11Hz),3.27(2H,q,J=7.5Hz),3.49(2H,a),3.7 9(1H,t,J=7.5Hz),4.08(2H,brs),7.20-7.35(5H,m),7.45(1 H,td,J=8,1.5Hz),7.49(1H,td,J=8,1.5Hz),7.74(1H,dd,J=8,1.5Hz),7.89(1H,td,J=8,1.5Hz),7.36(1 R spectrum ν (liq.) cm ⁻¹ :3360		64	н	Ph ₃ CN	3	mp,149-158°C Elemental analysis for C ₃₆ H ₃₇ ClN ₄ Calcd.%: C, 77.05; H, 6.85; N, 9.98
45 Mass spectrum m/z:394,396(M*,3:1)	40	65	Н	BnN	2	NMR spectrum & (ODCl ₃)ppm:1.20-1.50(3H,m),1.80(2H,q,J=7.5Hz),1.66(2H,d,J=11Hz),1.94(2H,t,J=11Hz), 2.88(2H,d,J=11Hz),3.27(2H,q,J=7.5Hz),3.49(2H,z),3.7 9(1H,t,J=7.5Hz),4.06(2H,brs),7.20-7.35(5H,m),7.45(1 H,td,J=8,1.5Hz),7.49(1H,td,J=8,1.5Hz),7.74(1H,dd,J=8,1.5Hz),7.89(1H,dd,J=8,1.5Hz)

	Reference	В	w	m	Physical properties
	example				(Recrystallization solvent)
5					colorless crystals (AcOEt-iso-Pr ₂ O)
			}		mp,167−167.5°C
	66	н	СН	0	Elemental analysis for C ₁₉ H ₂₅ ClN ₄ O ₂
10			{	1	Calcd.%: C, 60.55; H, 6.69; N, 14.87
					Found%: C, 60.47; H, 6.83; N, 14.81
					colorless crystals (iso-Pr ₂ O)
15	1				mp,154-155.5℃
15	67	CI	СН	2	Elemental analysis for C ₂₁ H ₂₈ Cl ₂ N ₄ O ₂
				1	Calcd.%: C, 57.40; H, 6.42; N, 12.75
					Found%: C, 57.31; H, 6.37; N, 12.69
20					coloriess crystals (iso-Pr ₂ 0)
					mp,129-129.5°C
	68	Me	CH	2	Elemental analysis for C ₂₂ H ₃₁ ClN ₄ O ₂
25					Calcd.%: C, 63.07; H, 7.46; N, 13.37
					Found%: C, 63.02; H, 7.56; N, 13.33
					colorless crystals (iso-Pr ₂ 0)
30	1				mp,140.5-141°C
	69	MeO	CH	2	Elemental analysis for C ₂₂ H ₃₁ ClN ₄ O ₃
					Calcd.%: C, 60.75; H, 7.18; N, 12.88
35					Found%: C, 60.61; H, 7.17; N, 12.81
					brown liquid
					NMR spectrum δ (CDCl ₃)ppm:1.14(2H,qd,J=12,3Hz),1.40-
					1.48(11H,m),1.50-1.70(5H,m),2.67(2H,t,J=12Hz),3.40(2H,t,
40	70	н	N	2	J=7.5Hz),4.07(3H,brs),7.39(1H,dd,J=8.5,4.5Hz),8.29(1H,dd
					4.5,2Hz),8.91(1H,dd,J=4.5,2Hz)=در
					IR spectrum ν (liq.) cm ⁻¹ :3344,2928,1694
45					Mass spectrum m/z:405,407(M+,3:1)

R³ NH

	r	1	1	
	Reference	R ²	R³	Physical properties
	example			(Recrystallization solvent)
5				colorless crystals (AcOEt-iso-Pr ₂ O)
				mp,115.5~116°C
	71	CI	BocN	Elemental analysis for C ₂₁ H ₂₉ ClN ₄ O ₂
10			<u> </u>	Calcd.%: C, 62.29; H, 7.22; N, 13.84
				Found%: C, 61.99; H, 7.28; N, 13.73
				colorless crystals (iso-Pr ₂ 0)
15		}	_	mp,132.5−134.5°C
	72	Me	BocN	Elemental analysis for C ₂₂ H ₃₂ N ₄ O ₂
			$\overline{}$	Calcd.%: C, 68.72; H, 8.39; N, 14.57
20				Found%: C, 68.65; H, 8.65; N, 14.48
				colorless prisms
				(iso-Pr ₂ O-n-Heptane)
25				mp,108-110℃
	73	CI	N Boc	Elemental analysis for C ₂₁ H ₂₈ ClN ₄ O ₂
			800	Caled.%: C, 62.29; H, 7.22; N, 13.84
				Found%: C. 62.18; H. 7.42; N. 13.81
30				colorless crystals (iso-Pr ₂ 0)
			^	mp,104–106°C
	74	CI		Elemental analysis for C ₂₁ H ₂₈ CIN ₄ O ₂
35			BocN	Calcd.%: C, 62.29; H, 7.22; N, 13.84
				Found%: C, 62.11; H, 7.35; N, 13.79
				colorless prisms (AcOEt-iso-Pr ₂ O)
40		75 CI	Boch	mp,128-128.5℃
	75			Elemental analysis for C ₂₀ H ₂₈ ClN ₅ O ₂
			∨ N~	Calcd.%: C, 59.18; H, 6.95; N, 17.25
45				Found%: C, 59.16; H, 6.84; N, 17.15

Reference	R ²	R ³	Physical properties
example			(Recrystallization solvent)
			green liquid
			NMR spectrum δ (CDCl ₃)ppm:1.47(9H,s),1.78
			2H,q,J=6Hz),2.69(1H,brs),2.99(1H,brs),3.30-
76	Ci		40(1H,m),3.50-3.55(1H,m),3.55-3.70(2H,m),3.
/0	G	Boch	5-4.05(3H,m),4.27(2H,brs),7.40-7.50(2H,m),7.
			0(1H,d,J=7.5Hz),7.90(1H,d,J=7.5Hz)
			IR spectrum v (liq.) cm ⁻¹ :3356,1696
			Mass spectrum m/z:406,408(M*,3:1)
			brown liquid
		BocHN	NMR spectrum & (GDCl ₂)ppm:1.40-1.55(2H,n
			,1.46(9H,s),2.00-2.05(2H,m),2.15-2.25(2H,m),
77	CI		45(2H,t,J=5.5Hz),2.80-2.90(2H,m),3.35(2H,t,c
"	G		5.5Hz),3.53(1H,brs),4.34(1H,brs),4.49(1H,brs)
			.40-7.50(2H,m),7.85-7.90(2H,m)
			IR spectrum v (liq.) cm ⁻¹ :3356,1694
			Mass spectrum m/z:419,421(M+,3:1)
			green liquid
			NMR spectrum δ (CDCl ₃)ppm:1.40-1.60(2H,r
78 Me			,1.46(9H,s),2.00-2.10(2H,m),2.10-2.25(2H,m),
		BocHN.	46(2H,t,J=5.5Hz),2.64(3H,a),2.85-2.90(2H,m)
	Me		.25(2H,t,J=5.5Hz),3.54(1H,brs),4.13(2H,brs),4
		\	9(1H,brs),7.39(1H,t,J=8.5Hz),7.44(1H,t,J=8.5
			z),7.89(1H,d,J=8.5Hz),7.91(1H,d,J=8.5Hz)
			IR spectrum v (liq.) cm ⁻¹ :3352,1704
			Mass spectrum m/z:399(M*)

	Reference	R ³	m	Physical properties
	example		L	(Recrystallization solvent)
	79	N Boc	2	coloriess plates (AcOEt-iso-Pr ₂ O) mp,104-105°C lelemental analysis for C ₂₀ H ₂₇ CIN ₁ O ₂ Calcd.S: O, 61.45; H, 6.96; N, 14.33 FoundS: O, 61.49; H, 6.81; N, 14.35 Specific rotation [α ₁ h ²⁰ : -20.9° (c=0.1, DMSO)
	80	\$\tag{\tag{\tag{\tag{\tag{\tag{\tag{	2	coloriess crystals (iso-Pr ₂ O) mp.96.5-99°C Elemental analysis for C ₁₈ H ₂₂ CIN ₄ O ₂ Calcd.%: C, 59.58; H, 6.39; N, 15.44 Found%: C, 59.30; H, 6.67; N, 15.30
,	81	но	2	coloriess crystals (AcOEt) mp.126-126°C Elemental analysis for C ₁₆ H ₃₁ CIN ₄ O Calcd.S: C, 59.90; H, 6.80; N, 17.46 FoundS: C, 59.71; H, 6.87; N, 17.32
5	82		2	yellowish brown liquid NMR spectrum δ (DCCl ₂)ppm:2.49(2H,t,J=5Hz),2.50 -2.80(4H,m),3.30-3.40(2H,m),3.75-3.85(4H,m),4.39(1 H,brs),4.50(2H,brs),7.44(1H,td,J=8.5,1Hz),7.48(1H,td,J=8.5,1Hz),7.91(1H,dd,J=8.5,1Hz),9.91(1H,d
,	83		3	yellowish brown liquid NMR spectrum & (CDCL),ppm:1.89(2H,quin,,J=8Hz),2. 45-2.60(4H,m),2.63(2H,t,J=6Hz),3.30(2H,t,J=6Hz),3. 78(4H,t,J=4.5Hz),4.50(3H,brs),7.44(1H,td,J=7.5,1Hz),7.47(1H,td,J=7.5,1Hz),7.83(1H,dd,J=7.5,1Hz),7.90(1H,dd,J=7.5,1Hz)) Right spectrum \(\nu\) (liq.) cm ⁻¹ :3344 Mass spectrum \(\nu\) (liq.) cm ⁻¹ :332(2(M', 3:1))

R³ NH NI

Reference R ³ example		Physical properties	
84		greenish brown liquid NMR spectrum & (CDCl ₂)ppm:1.45-1.80(2H,m),1.80-1.7((4H,m),2.35-2.60(4H,m),2.39(2H,t,J=5Hz),3.37(2H,t,J=5Hz),3.37(2H,t,J=5Hz),3.47(1H,td,J=7,1Hz),7.47(1Htd,J=7,1Hz),7.87(1H,dd,J=7,1Hz),7.84(1H,dd,J=7,1Hz) IR spectrum & (iiq.) cm ⁻¹ :3432,3340 Mass spectrum m/z:304,306(M*,3:1)	
85	⟨\n_	dark brown liquid NMR spectrum δ (CDCl ₃)ppm:1.80-1.90(4H,m),2.57(2H,t) J=5.5Hz),2.60-2.70(4H,m),3.40(2H,t,J=5.5Hz),4.27(3H,brs),7.43(1H,td,J=7.5.2Hz),7.46(1H,td,J=7.5.2Hz),7.87(1H,dd,J=7.5.2Hz),7.93(1H,dd,J=7.5.2Hz) IR spectrum ν (liq.) cm ⁻¹ :3436,3348 Mass spectrum m/z:290,292(M*,3:1)	

	Reference		Physical properties
	example		(Recrystallization solvent)
5		BocN	colorless crystals (iso-Pr ₂ 0)
			mp,130.5-131.5°C
	86	NH ₂	Elemental analysis for C ₂₁ H ₃₃ ClN ₄ O ₂
10			Calcd.%: C, 61.67; H, 8.13; N, 13.70
		√√N∕CI	Found%: C, 61.52; H, 8.29; N, 13.65
			colorless crystals
15	87	BocN	(CICH ₂ CH ₂ CI-iso-Pr ₂ O)
		V NH	mp.141.5~142.5℃
		NH ₂	Elemental analysis for C ₂₀ H ₃₁ ClN ₄ O ₂
20		(N C	Calcd.%: C, 60.82; H, 7.91; N, 14.19
		H CI	Found%: C, 60.63; H, 7.60; N, 14.03
25		BocN	gray orystals (AcOEt)
	88		mp,168-169°C
		NH NH ₂	Elemental analysis for C ₁₉ H ₂₇ ClN ₄ O ₂ S
			Calcd.%: C, 55.53; H, 6.62; N, 13.63
30		N CI	Found%: C, 55.54; H, 6.87; N, 13.63

Example 1

35 4-Chloro-1-[2-(N-triphenylmethyl-4-piperidyl)ethyl]-1H-imidazo[4,5-c]-quinoline

[0091] A solution of 19.9 g of 3-emino-2-chloro-4-[2-(N-triphenylmethyl-4-biperidyl)-ethyleminolquinoline, 24.1 m lof ethyl orthoformate and 0.68 g of p-toluenesulfonic acid monohydrate in 200 ml of toluene was refluxed for 8 hours. After cooling, the precipitated crystals were collected by filtration, and washed with disopropyl ether to give 16.9 g of coloriess crystals. Recrystallization from a mixture of methanol and tetrahydrofuran gave coloriess crystals naving the melting point of from 29 to 12.4 pc? (decomposition).

Elemental analysis for C ₃₆ H ₃₃ CIN ₄						
Calculated %	C, 77.61;	H, 5.97;	N, 10.06			
Found %	C, 77.50;	H, 5.98;	N, 9.95			

Example 2

4-Chloro-2-trifluoromethyl-1-[2-(N-triphenylmethyl-4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline

[0082] To a solution of 2.50 g of 3-amino-2-chloro-4-[2-(N-triphenylmethyl-4-piperidyl)ethylaminolquinoline and 0.76 ml of triethylamine in 60 ml of dried tetrahydrofluran, a solution of 0.63 ml of trifluoroacetic anhydride in 40 ml of dried tetrahydrofluran was addeed dropwise under ice-cooling, and the mixture was stirred at room temperature for 2-hours. The solvent of the reaction mixture was evaporated, and the residue was added with water and saturated aqueous sodium hydrogenocarbonate solution, and extracted with eithyl actate. The extract was washed successively with water and saturated brine, and dried, and then the solvent was evaporated. A solution of 3.03 g of the resulting pale yellow soil and 0.30 of of-follonessificine additional virtuation in 0.01 ml of tollone was refluxed for 20 hours. After the reactions

the solvent was evaporated, and the residue was added with methanol and acetone. The precipitated crystals were collected by filtration to give 1.79 g of colorless crystals.

NMR spectrum 8 (DMSO-d₆)ppm: 1.35-1.55(8H,m),1.59(2H,q,J=11Hz),1.77(2H,d,J=11Hz),1.80-1.90(2H,m).2.98(2H, brs),4.75(2H;L,J=8.8Hz),7.77(8H;L,J=88Hz),7.30(8H;L,J=8Hz),7.41(8H,brs),7.84(1H,Id,J=7.5,2Hz),7.87(1H,Id,J=7.5,2

Example 3

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tert-Butyl 4-[2-(4-methyl-2 -phenyl-1H-imidazo[4,5-c]quinolin-1-yl)ethyl]-1-piperidinecarboxylate

[0033] A solution of 0.65 g of tent-butyl 4/2-{(3-amino-2-methylquinolin-4y))amino-jethyl1-piperidinecarboylate, 0.29 g of benzaldehyde and 0.08 g of 2.3-dichloro-5, 6-dicyano-1, 4-benzoquinone in 5 mil of letrahydrofuran was stirred at room temperature for 3 days. The reaction mixture was added with saturated aqueous sodfum hydrogencarbonate solution and extracted with ethyl acetate. The extract was washed successively with saturated aqueous sodfum hydrogencarbonate solution and saturated brine, and dried, and the solvent was evaporated to give a reddsh brown liquid. The resulting liquid was purified by silica gel column chromatography using ethyl acetate - n-heptane (1:1) as an eluting solvent, and washed with dilsopropyl ether to give 0.55 g of a colorless solid. Recrystallization from dilsopropyl ether gave colorless crystals having the melling point of from 146 to 146.8°C.

Elemental analysis for C ₂₉ H ₃₄ N ₄ O ₂							
Calculated %							
Found %	C, 73.95;	H, 7.54;	N, 11.84				

25 [0094] In accordance with the methods of Examples 1 through 3, the compounds of Examples 4 through 72 were obtained.

Example	R1	В	m	Physical properties (Recrystallization solvent)
4	Н	Н	1	colorless crystals (MeOH) mp.232-239°C (decomposition) Elemental analysis for C ₅₅ H ₃₁ ClN ₄ Calcd.%: C, 77.40; H, 5.75; N, 10.32 Found%: C, 77.35; H, 5.79; N, 10.19
5	Ph	Н	1	pale yellow crystals (AcOEt) mp,165-168°C (decomposition) Elemental analysis for C ₄₁ H ₃₆ ClN ₄ Calcd.%: C, 79.53; H, 5.70; N, 9.05 Found%: G, 79.29; H, 5.74; N, 9.05
6	Н	CI	2	colorless crystals (MeOH) mp,266-268°C (decomposition) Elemental analysis for C ₉₆ H ₃₂ Cl ₂ N ₄ Calcd,%: C, 73.09; H, 5.45; N, 9.47 Found%: C, 73.15; H, 5.54; N, 9.41

(continued)

Example	R1	В	m	Physical properties (Recrystallization solvent)
7	Ph	Н	2	pale yellow crystals (CH ₂ Cl ₂ :EIOH) mp,246.5-249°C Elemental analysis for C ₄₂ H ₃₇ ClN ₄ Calcd.%: C, 79.66; H, 5.89; N, 8.85 Found%: C, 79.55; H, 6.12; N, 8.71
8	Ph	н	3	colorless crystals (AcOEI) mp,227.5-231°C (decomposition) Elemental analysis for C ₄₃ H ₃₆ ClN ₄ -1/4H ₂ O Calcd,%: C, 79.24; H, 6.11; N, 8.60 Found%: C, 79.26; H, 6.09; N, 8.55

R^-N (CH₂)_m

25	Example	R1	В	RA.	m	Physical properties (Recrystallization solvent)
	9	Н	Н	Bn	2	colorless crystals (AcOEt) mp,124.5-125°C Elemental analysis for C ₂₄ H ₂₅ ClN ₄ Calcd.%: C, 71.19; H, 6.22; N, 13.84 Found%: C, 71.22; H, 5.97; N, 13.79
30	10	Ph	Н	Boc	0	colorless crystals (AcOEt-MeOH) mp,250-255°C (decomposition) Elemental analysis for $C_{26}H_{27}ClN_4O_2$ Calcd.%: C, 67.45; H, 5.88; N, 12.10 Found%: C, 67.42; H, 5.88; N, 12.02
35	11	Н	Н	Boc	2	colorless crystals (AcOEt) mp,188-189°C Elemental analysis for C ₂₂ H ₂₇ ClN ₄ O ₂ Calcd,%: C, 63.68; H, 6.56; N, 13.50 Found%: C, 63.45; H, 6.60: N, 13.40
	12	Ph	CI	Boc	2	colorless crystals (AcOEt) mp,192-193°C Elemental analysis for \$C_{28}H_{30}Cl_2N_4C_2 Calcd.%: C, 64.00; H, 5.75; N, 10.66 Found%: C, 64.04; H, 5.59; N, 10.61
40	13	Ph	Me	Boc	2	colorless crystals (AcOEt) mp,182.5-183.5°C Elemental analysis for C ₂₉ H ₃₃ ClN ₄ O ₂ Calcd.%: C, 68.97; H, 6.59; N, 11.09 Found%: C, 68.91; H, 6.41; N, 11.06

R³

Example	В	R³	w	Physical properties (Recrystallization solvent)
	-		-	colorless crystals (AcOEt)
			1	mp.188.5~189.5°C
14	MeO	BocN	СН	Elemental analysis for C ₂₈ H ₃₃ ClN ₄ O ₃
14	meo		Un	Calcd.%: C, 66.85; H, 6.38; N, 10.75
			1	Found%: C, 66,70; H, 6,42; N, 10,70
	-		├	
			1	colorless crystals (MeOH)
		BocN	l	mp,225.5~227.5°C(decomposition)
15	н		N Elemental analysis for C ₂₇ H ₃₀ ClN ₅ C	
			1	Calcd.%: C, 65.91; H, 6.15; N, 14.23
			L	Found%: C, 65.85; H, 6.21; N, 14.21
		Book	СН	colorless crystals(AcOEt-n-Heptane)
	н			mp,159-161°C
16				Elemental analysis for C ₂₈ H ₃₁ ClN ₄ O ₂
				Calcd.%: C, 68.49; H, 6.36; N, 11.41
				Found%: C, 68.36; H, 6.27; N, 11.37
				colorless crystals (AcOEt-iso-Pr ₂ O)
		\sim	СН	mp,154.5-158°C
17	н	N Boc		Elemental analysis for C ₂₈ H ₃₁ ClN ₄ O ₂
				Calcd.%: C, 68.49; H, 6.36; N, 11.41
				Found%: C, 68.59; H, 6.15; N, 11.38
				colorless crystals (AcOEt)
		Bock		mp,166.5−167.5°C
18	н		CH	Elemental analysis for C ₂₈ H ₃₁ ClN ₄ O ₂
				Calcd.%: C, 68.49; H, 6.36; N, 11.41
			1	Found%: C, 68.50; H, 6.43; N, 11.32

R3 1

Example	R²	R³	Physical properties
			(Recrystallization solvent)
			colorless fine needles(AcOEt)
		_	mp,186.5−187.5°C
19	Cl	BocN	Elemental analysis for C ₂₇ H ₃₀ ClN ₅ O ₂
		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Calcd.%: C, 65.91; H, 6.15; N, 14.23
			Found%: C, 65.97; H, 6.31; N, 14.18
			colorless crystals (MeOH)
		_	mp,195.5~196.5°C
20	CI	Bock	Elemental analysis for C ₂₇ H ₂₉ ClN ₄ O ₃
			Calcd.%: C, 65.78; H, 5.93; N, 11.36
			Found%: C, 65.73; H, 5.86; N, 11.38
	СІ	BocHN	colorless crystals (AcOEt-iso-Pr ₂ O)
			mp,191.5−192°C
21			Elemental analysis for C ₂₈ H ₃₂ CIN ₅ O ₂
			Calcd.%: C, 66.46; H, 6.37; N, 13.84
			Found%: C, 66.42; H, 6.33; N, 13.69
		Me BocHN N	colorless crystals (AcOEt-iso-Pr ₂ O)
	Me		mp,164.5−165℃
22			Elemental analysis for C ₂₉ H ₃₅ N ₅ O ₂
			Calcd.%: C, 71.72; H, 7.26; N, 14.42
			Found%: C, 71.40; H, 7.24; N, 14.28

	Example	R¹	R³	m	Physical properties (Recrystallization solvent)
5	23	Ph	8	2	colorless crystals (AcOEt-iso-Pr ₂ O) mp,185-188°C Elemental analysis for C ₂₈ H ₂₈ CiN ₄ O ₂ Calcd.3: C, 86.88; H, 5.81; N, 12.48
					Found%: C, 66.59; H, 5.63; N, 12.45
15	24	Ph	но	2	colorless crystals (iso-PrOH) mp,184-170°C Elemental analysis for C ₂₂ H ₂₂ CIN ₄ O Calcd.3: C, 67.89; H, 5.70; N, 13.77 Found%: C, 67.82; H, 5.71; N, 13.63
<i>25</i> <i>30</i>	25	Ph		2	pale yellowish brown crystals (AcOEt) mp,182–183°C Elemental analysis for CzzHzrCIN ₄ O -1/4H ₂ O Calcd.%: C, 66.49; H, 5.45; N, 14.10 Found%: C, 66.26; H, 5.50; N, 14.03
35	26	Н		3	pale brown crystals (AcOEt) mp,130.5–131.5°C Elemental analysis for C ₁₇ H ₁₉ CIN ₄ O Calcd.%: C, 61.72; H, 5.79; N, 16.94 Found%: C, 61.72; H, 5.76; N, 16.90
45	27	Ph		3	pale brown crystals (MeOH) mp,183.5–184.5°C Elemental analysis for C ₂₂ H ₂₂ CIN ₄ O Caled.5: C, 67.89; H, 5.70; N, 13.77 Found%: C, 67.91; H, 5.66; N, 13.80

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	Example	RI	R³	m	Physical properties (Recrystallization solvent)
10	28	H		2	pale brown crystals (iso-Pr ₂ 0) mp.105-105.5°C Elemental analysis for C ₁₇ H ₁₈ ClN ₄ Calcd.5: C, 64.86; H, 8.08; N, 17.80 Found's: C, 64.83; H, 6.11; N, 17.72 pale brown crystals (MeOH) mp.226-227°C Elemental analysis for C ₂₂ H ₂₂ ClN ₄ Calcd.5: C, 70.67; H, 5.93; N, 14.33
20					Found%: C, 70.44; H, 5.96; N, 14.29
25	30	н	\(\frac{1}{2}\)	2	brown crystals NMR spectrum & (CDCl ₃)ppm:1.80-1.90(4H,m),2.58-2.76(4H,m),3.14-3.22(2H,m),4.78-4.91(2 H,m),7.68(1H,t,J=6.5Hz),7.72(1H,t,J=6.5Hz),8.1 3(1H,s),8.22(2H,d,J=6.5Hz) Mass spectrum m/z:300,302(M*,3:1)
35	31	Ph	⟨\n_	2	pale brown crystals (MeOH) mp,191–192°C Elemental analysis for C ₂₂ H ₂₁ ClN ₄ Calcd.%: C, 70.11; H, 5.82; N, 14.87 Found%: C, 70.00; H, 5.65; N, 14.86

	<u> </u>	
Example		Physical properties
Lampio		(Recrystallization solvent)
		colorless amorphous solid
		NMR spectrum δ (DMSO-d _B)ppm:0.99(3H,brs),
		32(3H,brs),1.68(2H,brs),2.13(1H,brs),2.49(9H,s),
	N Ph	.62-4.72(2H,m),7.60-7.67(3H,m),7.74-7.82(4H,n
32	Boc	,8.13(1H,dd,J=8,1.5Hz),8.42(1H,d,J=8Hz)
		IR spectrum ν (KBr)cm ⁻¹ :1690
	N, CI	Mass spectrum m/z:476,478(M+,3:1)
		Specific rotation
		[α] _p ²⁰ : -60.2° (c=0.1, DMSO)
	BocN	coloriess crystals (AcOEt)
	Ph Ph	mp,215-218°C (decomposition)
33		Elemental analysis for C ₂₈ H ₃₅ ClN ₄ O ₂
	I TYY	Calcd.%: C, 67.93; H, 7.13; N, 11.32
	N CI	Found%: C. 67.70; H, 7.17; N, 11.23
	BocN	colorless crystals (MeOH-iso-PrOH)
	Ph	mp,185-188°C
34		Elemental analysis for C ₂₇ H ₃₃ ClN ₄ O ₂
		Calcd.%: C, 67.42; H, 6.91; N, 11.65
	√ N CI	Found%: C, 67.31; H, 6.66; N, 11.57
	BocN	brown crystals (AcOEt)
	Ph Ph	mp,199-200°C
35		Elemental analysis for C ₂₈ H ₂₉ ClN ₄ O ₂ S
		Calcd.%: C, 62.83; H, 5.88; N, 11.27
	W ∩CI	Found%: C, 62.74; H, 5.83; N, 11.16

		Physical properties
Example	R ^t	(Recrystallization solvent)
		pale brown crystals (iso-PrOH)
ļ		mp,202-203°C
36	Me	Elemental analysis for C ₂₂ H ₂₉ CIN ₄ O ₂
-		Calcd.%: C, 64.40; H, 6.81; N, 13.06
1		Found%: C, 64.39; H, 7.04; N, 12.95
		colorless crystals (AcOEt-iso-Pr ₂ O)
		mp,159.5-160.5°C
37	n-Bu	Elemental analysis for C ₂₆ H ₂₅ ClN ₄ O ₂
1		Caled.%: C, 66.30; H, 7.49; N, 11.89
		Found%: C, 66.16; H, 7.53; N, 11.82
	a	colorless crystals (iso-PrOH)
		mp,174-175℃
38		Elemental analysis for C ₂₈ H ₃₇ CIN ₄ O ₂ • 1/4H ₂ O
1		Calcd.%: C, 67.05; H, 7.54; N, 11.17
1		Found%: C, 67.08; H, 7.47; N, 10.92
		colorless crystals (AcOEt-iso-Pr ₂ O)
		mp,165-166.5°C
39	Bn	Elemental analysis for C ₂₉ H ₃₃ ClN ₄ O ₂
1		Calcd.%: C, 68.97; H, 6.59; N, 11.09
1		Found%: C, 68.93; H, 6.72; N, 10.99
		coloriess crystals (AcOEt)
1	~0	mp,219-220.5℃ (decomposition)
40		Elemental analysis for C ₃₀ H ₃₃ CIN ₄ O ₂ ·1/4H ₂ O
1		Calcd.%: C, 69.08; H, 6.47; N, 10.74
		Found%: C, 69.25; H, 6.41; N, 10.69

BocN

		Physical properties
Example	R¹	(Recrystallization solvent)
		coloriess crystals (MeOH)
	, Me	mp,137~142℃
41		Elemental analysis for C ₂₉ H ₂₃ ClN ₄ O ₂ ·1/2H ₂ O
		Calcd.%: C, 67.76; H, 6.67; N, 10.90
1		Found%: C, 67.82; H, 6.49; N, 10.92
		colorless crystals (MeOH)
1	OMe	mp,153.5~157°C
42		Elemental analysis for C ₂₉ H ₃₃ ClN ₄ O ₃
		Calcd.%: C, 66.85; H, 6.38; N, 10.75
Í		Found%: C, 66.84; H, 6.54; N, 10.78
	"	colorless crystals (AcOEt)
1		mp,160-161°C
43		Elemental analysis for C ₂₈ H ₃₀ CIFN ₄ O ₂ ·1/8H ₂ O
Ì		Calcd.%: C, 65.78; H, 5.96; N, 10.96
		Found%: C, 65.57; H, 5.67; N, 10.94
		coloriess fine needles
1		(AcOEt-n-Heptane)
44		mp,180−182°C
**		Elemental analysis for C ₂₈ H ₃₀ CIFN ₄ O ₂
		Caled.%: C, 66.07; H, 5.94; N, 11.01
		Found%: C, 66.10; H, 5.71; N, 11.06
		colorless crystals (AcOEt-iso-Pr ₂ 0)
	✓ F	mp,126−129.5°C
45	Ü	Elemental analysis for C ₂₈ H ₃₀ CIFN ₄ O ₂
		Calcd.%: C, 66.07; H, 5.94; N, 11.01
		Found%: C, 66.06; H, 5.76; N, 11.01

			Physical properties
	Example	R¹	(Recrystallization solvent)
5		-	colorless crystals (iso-PrOH)
		F. J	mp,199.5−200°C
	46		Elemental analysis for C23H27CIF4N4O2
10		F	Calcd.%: C, 59.74; H, 4.83; N, 9.95
		· F	Found%: C, 59.61; H, 4.89; N, 9.90
		E	colorless crystals (iso-PrOH)
15		F. J. F	mp,216.5−217.5°C
	47		Elemental analysis for C ₂₈ H ₂₆ ClF ₅ N ₄ O ₂
		/ ¥ F	Calcd.%: C, 57.89; H, 4.51; N, 9.64
20		F	Found%: C, 57.88; H, 4.56; N, 9.62
			colorless crystals (AcOEt)
			mp,199.5-200.5°C
25	48		Elemental analysis for C ₂₇ H ₃₀ ClN ₅ O ₂
			Calcd.%: C, 65.91; H, 6.15; N, 14.23
			Found%: C, 65.77; H, 5.99; N, 14.25
30			colorless prisms
			(AcOEt-n-Heptane)
	49		mp,182−183°C
35	75	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	Elemental analysis for C ₂₇ H ₃₀ ClN ₅ O ₂
			Caled.%: C, 65.91; H, 6.15; N, 14.23
			Found%: C, 65.95; H, 6.26; N, 14.24
40			colorless prisms(AcOEt)
			mp,213-214℃
1	50		Elemental analysis for C ₂₃ H ₃₀ ClN ₅ O ₂
45			Calcd.%: C, 65.91; H, 6.15; N, 14.23
			Found%: C, 65.87; H, 6.20; N, 14.23

BocN

Example	R ^t	Physical properties
Example		(Recrystallization solvent)
		colorless crystals (MeOH)
1		mp,179−186°C
51		Elemental analysis for C ₂₉ H ₃₃ ClN ₄ O ₂ S
		Calcd.%: C, 64.85; H, 6.19; N, 10.43
1		Found%: C, 64.82; H, 6.45; N, 10.37
		colorless crystals (iso-PrOH)
	CF₃	mp,203-203.5°C
52		Elemental analysis for C29H30ClF3N4O2
}		Calcd.%: C, 62.31; H, 5.41; N, 10.02
		Found%: C, 62.24; H, 5.42; N, 9.99
	Ph	colorless crystals (AcOEt)
1		mp,224-225°C
53		Elemental analysis for C ₃₄ H ₃₅ ClN ₄ O ₂
		Calcd.%: C, 72.01; H, 6.22; N, 9.88
		Found%: C, 72.02; H, 6.21; N, 9.92
		colorless crystals (iso-PrOH)
	◇ OPh	mp,197−198°C
54		Elemental analysis for C ₃₄ H ₃₅ CIN ₄ O ₃
İ		Calcd.%: C, 70.03; H, 6.05; N, 9.61
		Found%: C, 69.83; H, 6.08; N, 9.58
		colorless crystals (MeOH)
		mp,196.5−197°C
55		Elemental analysis for C ₂₈ H ₂₉ GIN ₄ O ₃
		Galed.%: C, 64.93; H, 6.08; N, 11.65
1		Found%: C, 64.83; H, 6.27; N, 11.69

Example	R¹	R²	Physical properties (Recrystallization solvent)
56	; >	Me	Pale yellow crystals (iso-PrOH) mp.185.5–186*℃ Elemental analysis for C ₂₇ H ₁₂ N ₄ O ₃ Calcd.5: C, 70.41; H, 7.00; N, 12.16 Foundk: C, 70.32; H, 7.19; N, 12.13
57	\$	CI	colorless crystals (MeOH) mp,151.5-153°C Elemental analysis for C ₂₈ H ₂₉ CIN ₂ O ₂ S Calcd.X: C, 62.83; H, 5.88; N, 11.27 FoundX: C, 62.77; H, 6.01; N, 11.24
58	\$	Me	pale yellow crystals (iso-PrOH) mp,181.5-182.5°C Elemental analysis for C ₂₇ H ₃₂ N ₄ O ₂ S Calcd.X: C, 68.04; H, 6.77; N, 11.75 Found%: C, 67.86; H, 6.99; N, 11.63
59	s 🏠	CI	colorless crystals (AcOEt) mp.197–198°C Elemental analysis for C ₂₂ H ₂₆ CIN ₆ O ₁ S Calcd.X: C, 60.29; H, 5.67; N, 14.06 Found%: C, 59.98; H, 5.54; N, 13.84
60	5 N	Me	colorless crystals (AcOEt-iso-Pr ₂ O) mp,191-193°C Elemental analysis for C ₂₈ H ₂₁ N ₂ O ₂ S Calcd.X: C, 65.38; H, 6.54; N, 14.66 FoundX: C, 65.34; H, 6.53; N, 14.43

		Physical properties
Example	R¹	(Recrystallization solvent)
		yellow amorphous solid
		NMR spectrum & (GDCl ₂)ppm:
		1.08-1.09(2H,m),1.30-1.40(1H,m),140-1.45 (2H,m)
		1.44(9H,s),1.82-1.90(2H,m),2.55-2.82(2H,m),3.05(3
61		H,s),4.00-4.10(2H,m),4.62(2H,t,J=7.5Hz),7.27-7.30(
		2H,m),7.61(1H,t,J=7Hz),7.67-7.71(3H,m),8.14(1H,d.
		J=7.5Hz).8.24(1H.d.J=7.5Hz)
		IR spectrum v (KBr)cm ⁻¹ :1692
		Mass spectrum m/z:488(M*)
	F	colorless crystals (AcOEt)
	F, J. F	mp,195-196°C
62	ĬŢ,	Elemental analysis for C22H22F5N4O2
		Calcd.%: C, 62.14; H, 5.21; N, 9.99
		Found%: C, 62.07; H, 5.25; N, 9.94
		pale yellow crystals (AcOEt)
		mp,199.5~200.5°C
63		Elemental analysis for C28H32N5O2
1 1		Calcd.%: C, 71.31; H, 7.05; N, 14.85
		Found%: C, 71.37; H, 7.14; N, 14.83
		colorless crystals (MeOH-iso-Pr ₂ O)
	CF ₃	mp,177.5−179°C
64		Elemental analysis for C ₃₀ H ₃₃ F ₃ N ₄ O ₂
	~	Calcd.%: C, 66.90; H, 6.18; N, 10.40
		Found%: C, 66.89; H, 6.08; N, 10.37
1		pale brown crystals (AcOEt)
	HN 🥎	mp,193-194℃
65	``` <u>}</u>	Elemental analysis for C ₂₇ H ₃₃ N ₅ O ₂
		Galed.%: C, 70.56; H, 7.24; N, 15.24
		Found%: C, 70.61; H, 7.16; N, 15.21

Example	R¹	R²	Physical properties (Recrystallization solvent)
			colorless crystals (EtOH)
66	HN	CI	mp,240-241°C (decomposition) Elemental analysis for C ₂₅ H ₂₆ CiN ₅ O ₂
00)—N	L G	Caled.%: C. 62.43: H. 6.08: N. 17.47
			Found%: C, 62.49; H, 6.02; N, 17.51
			colorless crystals (EtOH)
	HN	Ме	mp,228.5-230°C (decomposition)
67			Elemental analysis for C ₂₆ H ₃₂ N ₆ O ₂
			Calcd.%: C, 67.80; H, 7.00; N, 18.25
			Found%: C, 67.72; H, 6.93; N, 18.24
			brown amorphous solid
	MeN		NMR spectrum δ (CDCl ₃)ppm:1.10-1.20(2H,m),1.4
		Me	6(9H,s),1.40-1.60(3H,m),1.90-1.98(2H,m),2.60-2.70
			2H,m),3.04(3H,s),3.86(3H,s),4.05-4.15(2H,m),4.74(2
68			H,t,J=8Hz),6.30(1H,t,J=2.5Hz),6.52(1H,d,J=2.5Hz),6
	/		88(1H,s),7.60(1H,t,J=8Hz),7.67(1H,t,J=8Hz),8.16(1H
			d,J=8Hz),8.23(1H,d,J=8Hz)
			IR spectrum ν (KBr)cm ⁻¹ :1688
			Mass spectrum m/z:473(M*)

			_	
	Example	R¹	R²	Physical properties
			<u> </u>	(Recrystallization solvent)
5			ĺ	yellow amorphous solid
	1			NMR spectrum & (CDCl ₃)ppm:
	69	S	CI	1.05-1.15(2H,m),1.40-1.50(3H,m),1.45(9H,s),1.83-1.90(
	03		G	2H,m),2.32(3H,s),2.60-2.70(2H,m),4.00-4.10(2H,m),4.60
10	ł	/ Me		-4.65(2H,m),7.06(1H,d,J=5.5Hz),7.51(1H,d,J=5.5Hz),7.6
	ł		l	8-7.75(2H,m),8.16(1H,d,J=7.5Hz),8.24(1H,d,J=7.5Hz)
		Me	1	pale yellow crystals (EtOH)
		me	l	mp.192-193°C
15	70	s	CI	Elemental analysis for C ₂₇ H ₃₁ CIN ₄ O ₂ S·5/4H ₂ O
) — /	1	Calcd.%: C, 60.77; H, 6.33; N, 10.50
		/	}	Found%: C, 60.82; H, 6.08; N, 10.17
			 	yellow amorphous solid
20		1		NMR spectrum δ (CDCI ₂)ppm:
				1.02-1.08(2H,m),1.44(9H,s),1.44-1.50(3H,m),1.80-1.90(
	1		1	2H.m),2,31(3H.s),2,60-2,70(2H.m),3,05(3H.s),4,00-4,05(
	71	\$	Me	2H.m).4.59(2H.t.J=7.5Hz).7.06(1H.d.J=5.5Hz).7.49(1H.d.
25	''	Me	mic	J=5.5Hz),7.60-7.65(2H.m),8.14(1H.d.J=8Hz),8.23(1H.d.J
		, we	1	=8Hz)
				IR spectrum ν (KBr)cm ⁻¹ :1688
			1	1
30			-	Mass spectrum m/z:490(M*)
		Me		pale yellow crystals (AcOEt)
		l Ï	1	mp,141~142°C
	72	s	Me	Elemental analysis for C28H34N4O2S-1/4H2O
35)=/		Calcd.%: C, 67.92; H, 7.02; N, 11.31
		/		Found%: C, 67.86; H, 6.84; N, 11.25

tert-Butyl 4-[2-(4-chloro-2-hydroxy-1H-imidazo[4,5-c]quinolin-1-yl)-ethyl]-1-piperidinecarboxylate

[0095] To a solution of 0.60 g of tent-butyl 4-(2-(3-amino-2-chioro-4-quinoylamino)-ethyl-1-piperidinecarboxylate is and 0.44 g of triphosgene in 10 ml of 1,2-dichloroethane, 0.41 ml of triethylamine was added dropwise, and the mixture was siltred air room temperature for 1 hour. The reaction mixture was neutralized with saturated aqueous sodium hydrogenoarbonate solution, and extracted with 1,2-dichloroethane. The extract was washed with saturated brine, and dried, and the solvent was experted. The residue was washed with discoproyl either to give 0.57 g of coloriess crystals Recrystallization from 1,2-dichloroethane gave coloriess crystals having the melting point of from 222 to 223°C.

Elemental analysis for C ₂₂ H ₂₇ CIN ₄ O ₃							
Calculated % C, 61.32; H, 6.32; N, 13.00							
Found % C, 61.15; H, 6.34; N, 13.00							

tert-Butyl 4-[2-[4-chloro-2-(4-methanesulfinylphenyl)-1H-imidazo[4,5-c]-quinolin-1-yl]ethyl]-1-piperidinecarboxylate

5 [0096] To a suspension of 0.63 g of tert-butyl 4:[24.chloro-24.4-methythio-phenyl)-H-imidazo[4,5-c]quinolin-1-yl] etlyl]-1-pperidinecarboxylate in 18 mil of 1,4-dioxane, a solution of 0.38 g of sodium periodate in 6 mil of water was added dropwise, and the mixture was stirred at 50°C for 13 hours. The reaction solution was concentrated, and the residue was purified by silica get column chromatography using 1,2-dichloroethane - methanol (10:1) as an eluting solvent to give 0.47 g of a colorless solid. Recrystallization from a mixture of isopropanol and water gave colorless crystals having the mething opin of from first 30 to 186°C.

Elemental analysis for C ₂₉ H ₃₃ ClN ₄ O ₃ S · 1/4H ₂ O								
Calculated % C, 62.46; H, 6.06; N, 10.05								
Found %								

Example 75

tert-Butyl 4-[2-[4-chloro-2-(4-methanesulfonylphenyl)-1H-imidazo[4,5-c]-quinolin-1-yl]ethyl]-1-piperidinecarboxylate

[0097] To a solution of 0.40 g of tert-butyl 4:[2:[4-chloro-2:(4-methythiophenyl)-1H-imidazo[4.5 -clgulnoin-1-yl] ethyl-1-peindineaerboxylate in 20m tof 1.2-dichloroethane, 0.40 g of m-chloroperboxnoic acid was added portionwise little by little, and the mixture was stirred at room temperature for 1 hour. The reaction mixture was neutralized with 10% aqueous sodium hydroxide solution, and extracted with 1.2-dichloroethane. The extract was washed with saturated aqueous sodium hydrogenorabonate solution and dried, and then the solvent was evaporated. The residue was washed with a mixture of disopropyl ether and diethyl ether to give 0.42 g of colorless crystals. Recrystall zation from methanol cave colorless crystals having the mediting point of from 149 to 156°C.

Elemental analysis for C ₂₉ H ₃₃ ClN ₄ O ₄ S · 1/4H ₂ O							
Calculated % C, 60.72; H, 5.89; N, 9.77							
Found % C, 60.72; H, 5.81; N, 9.67							

Example 76

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4-Hydroxy-2-phenyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline

[0098] A solution of 871 mg of 4-chloro-2-phenyl-1-[2-(4-piperidylpethyl]-1H-imidazo(4,5-0-jquinoline and 2.5 ml of 6 N hydrochloric acid in 8 ml of 1,4-dioxane was refluxed for 3 hours. The reaction mixture was adjusted to pH 10 with 10%, acqueous sodium hydroxide solution, and added with potassium carbonate, and then extracted with 1.2-dichloroethane. The extract was dried, and the solvent was evaporated. The resulting residue was washed with ethyl acetate to give 522 mg of pale brown crystals. Recrystallization from methanol gave pale brown crystals having the metting point of from 242 5 to 244°C.

Elemental analysis for C ₂₃ H ₂₄ N ₄ O · 1/4H ₂ O						
Calculated % Found %	C, 73.28; C, 73.32;					

[0099] In accordance with the method of Example 76, the compounds of Examples 77 through 79 were obtained.

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	Example	В	R³	m	Physical properties (Recrystallization solvent)
5	77	CI	BnN	2	oolorless crystals (MeOH) mp,269-280°C (decomposition) Elemental analysis for C ₂₄ H ₂₅ CIN ₄ O Calcd.%: C, 68.48; H, 5.99; N, 13.31 Found%: C, 68.32; H, 8.07; N, 13.29
15	78	Н	HN	1	colorless crystals [hydrochloride] NMR spectrum & (DMSO-d ₀)ppm: 1.56(2H ₄ ,J ₌ 11.5H ₂),1.74(2H ₄ J,J ₌ 11.5H ₂),2.10-2.2 5(1H ₁ m),2.79(2H ₄ J,J ₌ 11.5H ₂),3.24(2H ₄ J,J ₌ 11.5H ₂),4.54(2H ₄ J,J ₌ 1.5H ₂),7.49(1H ₄ J,J ₌ 8H ₂),7.50(1H ₄ J,J ₌ 8H ₂),8.36(1H ₄ J,J ₌ 8H ₂
25 30	79	н	BnN	1	colorless crystals [hydrochloride] NMR spectrum δ (DMSO-d ₃)ppm: 1.65-1.85(4H,m),2.00-2.15(1H,m),2.84(2H,q,J=12H 2.).3.30(2H,d,J=12H2),4.18(2H,d,J=5H2),4.51(2H,d, J=7.5H2),7.27(1H,t,J=6.5H2),7.40-7.60(7H,m),7.97 (1H,d,J=8H2),8.31(1H,b.).10.83(1H,brs),11.58(1H,s) IR spectrum ν (KBr) cm ⁻¹ .3416.1672 Mass spectrum m/z:3720M*)

35 Example 80

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tert-Butyl 4-[2-(4-phenoxy-1H-imidazo[4,5-c]quinolin-1-yl)ethyl]-1-piperidinecarboxylate

[0100] A mixture of 4.46 g of tert-bulyl 4-12-(4-chlore-1H-imidaze)(4.5-clguinolin-1-y)bethyl-1-piperidinecarboxylate, 0.10 g of phenoi and 1.80 g of potassaium hydroxide was stirred at 1907° for 7 hours. The reaction mixture was adjusted to pH 10 with 10% aqueous sodium hydroxide solution, and extracted with ethyl acetate. The extract was washed successively with 10% aqueous sodium hydroxide solution and saturated brine, and dried, and then the solvent was evaporated to give a brown liquid. The resulting brown liquid was purified by sitilise gol column chromadgraphy using othyl acetate as an eluting solvent to give 3.59 g of a colorless solid. Recrystallization from a mixture of ethyl acetate and n-hoxane gave colorless crystals having the melting point of from 130.5 to 132.5°C.

Elemental analysis for C ₂₈ H ₃₂ N ₄ O ₃							
Calculated % C. 71.16; H, 6.83; N, 11.86							
Found % C, 71.10; H, 7.10; N, 11.69							

[0101] In accordance with the method of Example 80, the compounds of Examples 81 through 87 were obtained.

Example	R ¹	R³	R ⁸	Physical properties (Recrystallization solvent)
81	н	BnN	н	coloriess crystals (MeOH) mp.152.5-153.5°C Elemental analysis for C ₂₀ H ₃₀ N ₄ O Calcd.S: C, 77.89; H, 6.54; N, 12.11 FoundS: C, 78.00; H, 6.29; N, 12.05
82	н	AcN	н	oolorless orystals (AcOEt-iso-Pr ₂ O) mp.187-189.5°C Elemental analysis for C ₂₅ H ₂₈ N ₄ O ₂ Calcd.S. C. 72.44; H, 6.32; N, 13.52 FoundS: C, 72.35; H, 6.26; N, 13.42
83	н	AcN	F	ooloriess crystals (CH ₂ Cl ₂ -iso-Pr ₂ O) mp.206.5–208°C Elemental analysis for C ₂₃ H ₂₈ FN ₄ O ₂ ·1/8H ₂ O Calod.S. C, 69.07; H, 5.85; N, 12.89 FoundS: C, 69.11; H, 5.74; N, 12.85
84	Ph	AcN	н	colorless crystals (MeOH-iso-Pr ₂ O) mp.205-207.5°C Elemental analysis for C ₃₁ H ₃₀ N ₄ O ₂ -1/2H ₂ O Calcd,S: C, 74.53; H, 6.25; N, 11.21 Founds: C, 74.52; H, 6.37; N, 11.10

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Example	R ¹	R³	R [®]	Physical properties (Recrystallization solvent)
85	н	BocN	F	colorless crystals (AcOEt-n-Hexane) mp.133.5-135.5°C Elemental analysis for C ₂₈ H ₃₁ FN ₄ O ₂ Calcd.S: C, 68.55; H, 6.37; N, 11.42 Found%: O, 68.37; H, 6.47; N, 11.25
86	Ph	BocN	н	coloriess crystals (iso-PrOH) mp,207-208 [®] C Elemental analysis for C ₃₄ H ₃₆ N ₄ C ₃ Calcd.%: C, 74.43; H, 6.61; N, 10.21 Found%: C, 74.38; H, 6.68; N, 10.14
87	Н	\(\sigma_n\)	Н	pale purple crystals NMR spectrum 6 (DMSO-d_p)ppm: 1.64-1.72(4H,m).2.55-2.58(4H,m).2.88(2H,t,J=7 Hz).4.80(2H,t,J=7Hz).7.25-7.31(3H,m).7.45-7.4 9(2H,m).7.53-7.80(2H,m).7.72(1H,d,J=7Hz).8.29 (1H,d,J=7Hz).8.37(1H,s) Mass spectrum m/z:358(M*)

Example 88

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tert-Butyl 4-[2-(4-amino-1H-imidazo[4,5-c]quinolin-1-yl)ethyl]-1-piperidinecarboxylate

[0102] A mixture of 4.40 g of tert-buly 4/2-(4-phenoxy-11-limidazo(4.5-(-quinolin-1-y)bethyl1-siperidinecarboxyiate and 4.5 g of ammonium caetate was stirred at 140°C for 8 hours. The reaction mixture was added with vater, acquisted to pH 9 with 10% aqueous sodium hydroxide solution, and extracted with methylene chloride. The extract was washed with sarurated brine, and dried, and then the solvent was evaporated. The resulting residue was purified by alumina column chromatography using methylene chloride - methanol (100:1 to 20:1) as eliting solvents, and washed with diisopropyl ether to give 1.88 g of coloriess crystals. Recrystallization from ethyl acetate gave coloriess crystals having the melting point of from 198 to 19.85°C.

Elemental analysis for C ₂₂ H ₂₉ N ₅ O ₂							
Calculated % C, 66.81; H, 7.39; N, 17.71							
Found % C, 66.93; H, 7.48; N, 17.66							

[0103] In accordance with the method of Example 88, the compounds of Examples 89 through 92 were obtained.

Example	R³	Physical properties (Recrystallization solvent)
89	BnN	oolorless crystals (EtOH) mp,191.5–192°C Elemental analysis for C ₂₄ H ₂₇ N ₂ Calcd.S. C, 74.77; H, 7.06; N, 18.17 FoundS: C, 74.87; H, 7.18; N, 18.06
90	Acn	rounds: C, 17.5.7; N, 17.15; N, 18.00 colorless crystals (MeOH) mp,231.5–232.5°C Elemental analysis for C ₁₉ H ₂₉ N ₆ O Caled.5: C, 67.63; H, 6.87; N, 20.76 Founds: C, 67.46; H, 6.79; N, 20.63
91	EtO ₂ CN	colorless crystals (EtOH) mp.166–167°C Elemental analysis for C ₂₀ H ₂₅ N ₅ O ₂ Calod.5: C, 65.37; H, 6.86; N, 19.06 Found%: C, 65.52; H, 6.76; N, 18.83
92	Q-	pale yellow crystals [fumarate] (DMF-isc-Pr ₂ O) mp,195-197°C (decomposition) Elemental analysis for C1 ₁ H ₁₁ N ₁ · C ₄ H ₄ O ₄ · 5/4H ₄ O Calcd.X: C, 57.20; H, 6.12; N, 16.68 FoundX: C, 57.20; H, 6.23; N, 16.53

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tert-Butyl 4-[2-(4-dimethylamino-2-phenyl-1H-imidazo[4,5-c]quinolin-1-yl)-ethyl]-1-piperidinecarboxylate

[0104] A mixture of 0.89 g of tert-butyl 4-[2-(4-chloro-2-phenyl-1H-imidazo[4,5-6]-quinolin1-1ylpethyl]-1-piperidine-carboxylate and 7 ml of 50% aqueous dimethylamine solution was stirred in a sealed tube at 80°C of outer temperature for 2 hours. The reaction solution was added with water and extracted with only accelate. The extract was weahed successively with water and saturated brine, and dried, and the solvent was evaporated. The residue was washed successively with isopropanol and disopropyl either to give 0.52 g of colorless crystals. Recrystallization from isopropanol gave colorless crystals having the melting point of from 170.5 to 171.5°C.

Elemental analysis for C ₃₀ H ₃₇ N ₅ O ₂								
Calculated % C, 72.12; H, 7.46; N, 14.02								
Found %								

50 Example 94

tert-Butyl 4-[2-[4-(4-methylpiperazin-1-yl]-2-phenyl-1H-imidazo[4,5-c]-quinolin-1-yl]ethyl]-1-piperidinecarboxylate

[016] A mixture of 0.80 g of tert-buty 4-12-(4-chloro-2-phenyl-1H-imidaze-[4,5-clquinolin1-yhghty]t-1-pieridine-carboxylate and 1 ml of N-methylopierazine was stirred at 80°C for 6 hours. The reaction mixture was added with saturated aqueous sodium hydrogenerathorate solution and extracted with ethyl acetate. The extract was dried, and the solvent was evaporated. The residue was purified by alumina column chromatography using ethyl acetate - n-hectane (13.6 1.1) as eluting solvents, and washed with a mixture of disoporopy other and n-hepotane to dive 0.74 el

of colorless crystals. Recrystallization from ethyl acetate gave colorless needles having the melting point of from 140 to 141 °C.

Elemental analysis for C ₃₃ H ₄₂ N ₆ O ₂					
Calculated % C, 71.45; H, 7.63; N, 15.15					
Found %	C, 71.23;	H, 7.65;	N, 14.99		

[0106] In accordance with the methods of Examples 93 and 94, the compounds of Examples 95 through 102 were obtained.

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Example	R²	Physical properties
		(Recrystallization solvent)
- 1		colorless crystals (iso-PrOH)
ļ		mp,161-162°C
95	NHMe	Elemental analysis for C ₂₉ H ₃₅ N ₅ O ₂ ·1/2H ₂ O
		Calcd.%: C, 70.42; H, 7.34; N, 14.16
		Found%: C, 70.31; H, 7.23; N, 13.95
1		colorless crystals (iso-Pr ₂ O)
	^	mp,162-162.5°C
96	N Z	Elemental analysis for C ₃₁ H ₃₇ N ₅ O ₂ ·1/2H ₂ O
	Ĥ	Caled.%: C, 71.51; H, 7.36; N, 13.45
-		Found%: G, 71.73; H, 7.35; N, 13.09
		colorless needles (MeOH)
		mp,171-172℃
97		Elemental analysis for C ₃₃ H ₄₁ N ₅ O ₂
1		Calcd.%: C, 73.44; H, 7.66; N, 12.98
1		Found%: C, 73.44; H, 7.88; N, 12.93
1		colorless crystals (iso-PrOH)
i	\n\	mp,189-190°C
98	Ĩ	Elemental analysis for G ₃₂ H ₃₉ N ₅ O ₃
1	<u></u>	Calcd.%: C, 70.95; H, 7.26; N, 12.93
		Found%: C, 71.22; H, 7.47; N, 12.94
1		pale brown amorphous solid
-		NMR spectrum δ (CDCl ₃)ppm:
į.		0.99-1.06(2H,m),1.25-1.40(3H,m),1.43(9H,s),1.80-1.
i		90(2H,m),2.50-2.60(2H,m),3.95-4.05(2H,m),4.59(2H,
99	NHBn	,J=7.5Hz),4.96(2H,d,J=5.5Hz),8.11(1H,t,J=5.5Hz),7.2
}	1	4-7.28(1H,m),7.30-7.35(3H,m),7.48(2H,d,J=7.5Hz),7
		50-7.55(4H,m),7.60-7.65(2H,m),7.94-7.96(2H,m)
		IR spectrum ν (KBr) cm ⁻¹ :3436,1690
		Mass spectrum m/z:561(M*)

BooN

Example	R²	Physical properties
		pale yellow amorphous solid
		NMR spectrum δ (CDCl₃)ppm:
		1.00-1.08(2H,m),1.30-1.35(1H,m),1.38-1.42(2H,m),1.
	^	43(9H,s),1.83-1.90(2H,m),2.57(2H,brs),3.98(2H,brs),4
100	4 1 1	.61(2H,t,J=7.5Hz),4.99(2H,d,J=6Hz),7.33-7.35(1H,m),
		7.39(2H,d,J=6Hz),7.51-7.59(4H,m),7.64-7.67(2H,m),7
		.88-7.89(1H,m),7.96-7.97(1H,m),8.53(2H,d,J=6Hz)
		IR spectrum ν (KBr) cm ⁻¹ :3428,1692
		Mass spectrum m/z:562(M*)
		pale brown amorphous solid
Ì		NMR spectrum δ (CDCl ₂)ppm:
į) OMe	0.98-1.06(2H,m),1.25-1.40(3H,m),1.43(9H,s),1.80-1.
l		85(2H,m),2.50-2.60(2H,m),3.79(3H,s),3.90-4.00(2H,m
101),4.59(2H,t,J=7.5Hz),4.87(2H,d,J=5.5Hz),6.05(1H,brs)
101		,6.86(2H,d,J=8.5Hz),7.31(1H,t,J=7.5Hz),7.40(2H,d,J=
- 1		8.5Hz),7.51-7.60(4H,m),7.60-7.65(2H,m),7.94(2H,d,J
1		=8.5Hz)
		IR spectrum ν (KBr) cm ⁻¹ :3432,1692
		Mass spectrum m/z:591(M ⁺)
		colorless amorphous solid
		NMR spectrum δ (DMSO-d _e)ppm:
- 1		0.87(2H,q,J=5Hz),1.20-1.35(3H,m),1.36(9H,s),1.75(2
		H,q,J=7.5Hz),2.54(2H,t,J=12.5Hz),3.77(2H,d,J=12.5H
102		z),4.64(2H,t,J=7.5Hz),6.99(1H,t,J=8Hz),7.34(2H,t,J=8
	N N	Hz),7.44(1H,t,J=7.5Hz),7.56(1H,t,J=7.5Hz),7.60-7.67
	п .	(3H,m),7.76-7.82(2H,m),7.87(1H,d,J=7.5Hz),8.16(1H,
		d,J=7.5Hz),8.24(2H,d,J=8Hz),9.03(1H,s)
1		IR spectrum \(\nu\) (KBr) cm ⁻¹ :2932,1692
		Mass spectrum m/z:547(M ⁺)

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4-Amino-2-phenyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline trifluoroacetate

[0107] A mixture of 0.30 g of tert-butyl 4-[2-[4-(4-methoxybenzylamino)-2-phenyl-11-lmidazo[4,5-c]quinolin-1-yl] ethyl]-1-pperdimecarboxylate and 9 ml of trilluoroacetic acid was stirred at 65°C of outer temperature for 8 hours. The reaction solution was concentrated, and the residue was added with isopropane. The precipitated crystals were oclicated by filterition, and washed with diisopropyl either to give 0.31 g of pale yellow crystals. Recrystallization from a mixture of eithanol and isopropanel given colorises crystals having the meltitip goint of from 229 to 224°C.

Elemental analysis for C ₂₃ H ₂₅ N ₅ · 2CF ₃ CO ₂ H · H ₂ O					
Calculated % C, 52.51; H, 4.73; N, 11.34					
Found %	C, 52.61;	H, 4.45;	N, 11.61		

1-[2-(4-Chloro-2-phenyl-1H-imidazo [4,5-c]quinolin-1-yl)ethyl]-4-piperidinone

5 [0108] A mixture of 0.39 g of 11₂/4-d-blore-2-phenyl-1H-imidazo(A, 5-Gquinolin-1-y)lethyl|-4-d-ethylenedioxyp peridne and 4 mol concentrated sulfurie acid was stirred at room temperature for 30 minutes. The reaction mixture was poured into ice-water, adjusted to pH 11 with 10% aqueous sodium hydroxide solution, and extracted with othly acidate. The extract was washed with saturated aqueous sodium hydroxide solution, and dried, and then the solvent was evaporated to give 0.42 g of a colorless liquid. The resulting figuid was purified by alumina column chromatography using othly acotate - n-heptane (1:1) as an oluting solvent to give 0.32 g of colorless crystals. Recrystallization from isopropened awe colorless eneddles having the melting point of from 163 to 165°C.

Elemental analysis for C ₂₃ H ₂₁ ClN ₄ O					
Calculated % C, 68.23; H, 5.23; N, 13.84					
Found %	C, 68.26;	H, 5.31;	N, 13.78		

Example 105

20 1-[2-(4-Chloro-2-phenyl-1H-imidazo[4,5-c]quinolin-1-yl)ethyl]-4-piperidinone oxime

Elemental analysis for C ₂₃ H ₂₂ CIN ₅ O · 1/2H ₂ 0					
ĺ	Calculated %				
	Found %	C, 64.75;	H, 5.32;	N, 16.09	

Example 106

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tert-Butyl 4-[2-(2-phenyl-1H-imidazo[4.5-clquinolin-1-yl)ethyl]-1-piperidinecarboxylate

[0110] A suspension of 0.80 g of tert-butyl 4-[2-(4-chloro-2-phenyl-1H-imidazo-[4,5-c]quinolin-1-yllethyll-1-piperidinecarboxylate and 0.30 g of 5% palladium on carbon in 80 ml of methanol was catalytically hydrogenated at ordinary temperature under atmospheric pressure for 12 hours. After the reaction, the catalyst was filtered off, and the filtrate was concentrated. The residue was purified by silica gel column chromatography using ethyl acetato - n-heptane (1: 1 to 4:1) as eluting solvents and washed with diisopropyl ether to give 0.49 g of pale yellow crystals. Recrystallization from disporrow there gave coloriess crystals having the melting opinion from first 8 to 159°C.

Elemental analysis for C ₂₈ H ₃₂ N ₄ O ₂				
Calculated % C, 73.66; H, 7.06; N, 12.27 Found % C, 73.46; H, 7.21; N, 12.17				

[0111] In accordance with the method of Example 106, the compounds of Examples 107 through 109 were obtained.

Example	R³	m	Physical properties (Recrystallization solvent)
107	(MeOH) mp,258-2 1 Elementa C ₁₀ H ₁₈ N ₄ Calod.3:		coloriess crystals [hydrochloride] (MaCH) mp_258-261°C (decomposition) Elemental analysis for C ₁₆ H ₁₀ M ₂ ·22HCI-H ₂ O CaloXi: C, 53.79; H, 6.21; N, 15.68 Found%: C, 53.49; H, 6.14; N, 15.67
108	HN	2	colorless crystals [hydrochloride] (MeOH-ClCH ₂ CH ₂ (I) mp.220-233°C (decomposition) Elemental analysis for C ₁₇ H ₂₈ N ₄ -2HCl-1/2H ₂ O Calcd.S: C, 56.38; H, 6.40; N, 15.48 FoundS: C, 56.36; H, 6.18; N, 15.25
109	n-BuN	2	coloriess crystals [hydrochloride] (MeOH-iso-Pr ₂ O) mp.225-238°C (decomposition) Elemental analysis for C ₂₁ H ₂₀ N ₄ , 22HCl-1/8H ₂ O Galcd.S. C, 61.27; H, 7.41; N, 13.61 FoundS: C, 61.03; H, 7.44; N, 13.50

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4-Chloro-2-phenyl-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4.5-c]quinoline hydrochloride and fumarate

[0112] A mixture of 3.84 g of 4-chloro-2-phenyl-1-12-(N-triphenylmethyl-4-piperidy)lethyl]-11-limidaz(4.6-jcluinoline, 30 ml of methanol and 10 ml of trifluoroacetic acid was stirred at room temperature for 1 hour. The reaction mixture was concentrated, and the residue was washed successively with othyl acotate and diethyl other to give pale brown crystals (influoroacetals). The resulting crystals were added with ethyl acotate, and extracted with water. The aqueous layer was adjusted to pl-11 with 10% aqueous sodium hydroxide solution, and extracted with mature of 1.2-dichloroethane and methanol. The extract was washed with saturated brine, and dried, and then the solvent was evaporated to give 1.74 g of a colorless liquid. A part of the colorless liquid was convented into hydrochloride in a conventional method. Recrystallization from methanol gave colorless crystals having the metiting point of from 257 to 256°C (decomposition). In the same manner, fumarate was prepared in a conventional method. Recrystallization from methanol gave colorless crystals having the metiting point from 185.5 to 168.5°C (decomposition).

Hydrochloride:

0 [0113]

Elemental analysis for C ₂₃ H ₂₃ CIN ₄ · HCI · H ₂ O					
Calculated % C, 62.02; H, 5.88; N, 12.58 Found % C, 62.08; H, 5.77; N, 12.60					
Found %	C, 62.08;	H, 5.77;	N, 12.60		

Fumarate:

[0114]

Elemental analysis for C ₂₃ H ₂₃ ClN ₄ · C ₄ H ₄ O ₄ · H ₂ O					
Calculated % Found %	C, 61.77; C, 62.04;				

Example 111

4-Phenoxy-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4.5-c]quinoline trifluoroacetate

[0115] To a solution of 0.30 g of tert-buty14.[2-(4-phenoxy-1H-imidazo(14.5-)-quinolin-1-y)-leihyl]-1-pipordinicacrioxylate in 10 ml of methylene chloride, 1 ml of trifluoroscelic acid was added at room temperature, and the mixture was stirred for 1.5 hours. The reaction solution was concentrated. The resulting pale yellow solid was washed successively with isopropanol and dilospropyl either to give 0.38 g of colorless crystals. Recrystallization from a mixture of methylene chloride and eithanol gave colorless crystals having the melting point of from 211 to 216°C.

Elemental analysis for C23H24N4O · CF3CO2H · 1/8H2O				
Calculated %	C, 61.44;	H, 5.21;	N, 11.46	
Found %	C, 61.26;	H, 5.05;	N, 11.47	

os Example 112

20

4-Chloro-2-phenyl-1-[2-(1-piperazinyl)ethyl]-1H-imidazo[4,5-c]quinoline methanesulfonate

[0116] To a solution of 1.20 g of tert-butyl 4-[2-(4-chlore-2-phenyl-1H-limidaze-)4,5-c]quinoin-1-y)lethyl-1-piperazi-necarboxylate in 12 ml of 1,2-dichloroethane, 1.2 ml of methanesulfonic acid was added, and the mixture was stirred at room temperature for 5 minutes. The reaction mixture was added with isopropanol and ethanol, and the prepitiated crystals were collected by filtration to give 1.24 g of colorises crystals. Recrystallization from methanol gave coloriess crystals having the melting colori for for 26 to 26 of 20°C (docomosottion).

Elemental analysis for C ₂₂ H ₂₂ CIN ₅ · 2CH ₃ SO ₃ H					
Calculated % C, 49.35; H, 5.18; N, 11.99					
Found %	C, 49.60;	H, 5.11;	N, 12.16		

40 Example 113

55

4-Amino-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]quinoline hydrochloride

[0117] A mixture of 1.57 g of tert-butyl 4;[2;4]-aminor-1H-imidazo(4,5-lq)uinolin-1-y)lethyl-1-piperidinecarboxylate and 40 ml of ethyl acetate solution of hydrogen chloride was stirred at room temperature for 5 hours. The reaction mixture was added with water, adjusted to pH 10 with 10% aqueous sodium hydroxide solution, and extracted with methylene chloride. The extract was dried, and the solvent was evaporated. The resulting residue was washed with ethyl acetate to give 1.01 g of pelbe brown crystals. The resulting roystals were purified by alumina column chromatography using methylene chloride - methanol (40:1 to 20:1) as eluting solvents, and washed with disopropyl either to give coloriess crystals. Hydrochloride was prepared in a conventional method. Recrystallization from ethanol gave coloriess crystals having the melting point of from 243 to 244°C (decomposition).

Elemental analysis for C ₁₇ H ₂₁ N ₅ · HCl · 3/4H ₂ O				
Calculated %	C, 59.12;	H, 6.86;	N, 20.28	
Found %	C, 59.10;	H, 6.83;	N, 20.30	

[0118] In accordance with the methods of Examples 110 through 113, the compounds of Examples 114 through 186

were obtained.

HN (CH₂)m N

Example	R1	В	m	Physical properties (Recrystallization solvent)
114	Ph	н	0	colorless crystals (CICH ₂ CH ₂ CI-AcOEt) mp,253-256°C (decomposition) Elemental analysis for C ₂₁ H ₁₉ CIN ₄ Calcd.%: C, 69.51; H, 5.28; N, 15.44 Found%: C, 69.29; H, 5.19; N, 15.27
115	Н	Н	1	colorless crystals [hydrochloride] (McOH-EOH) mp,273-286*C (decomposition) Elemental analysis for C ₁₆ H ₁₇ ClN ₂ -2HCl Calcd, %: C, 51.42; H, 5.12; N, 14.99 Found%: C, 51.47; H, 5.08; N, 14.85
116	Ph	Н	1	colorless crystals [fumarato](MeOH) mp.288-271.5°C (decomposition) Elemental analysis for C ₂₉ H ₂₂ CIN ₄ -1/2C ₄ H ₂ Q ₄ -5/2H ₂ Q Calcd,%: C, 62.40; H, 5.67; N, 12.13 Found%: C, 62.52; H, 5.28; N, 12.15
117	Н	Н	2	colorless crystals [hydrochloride] (ETOH) mp_258-267*C (decomposition) Elemental analysis for C ₁₇ H ₁₆ ClN ₂ -HCl Calcd, %: C, 58, 13; H, 5.74; N, 15.95 Found%: C, 57.88; H, 5.46; N, 15.78
118	Н	СІ	2	colorless crystals [trifluoroacetate] (MeOH-iso-Pr _p O) mp_204-207.5°C Elemental analysis for C ₁₇ H, gC ₂ N ₄ ·CF ₂ CO ₂ H-1/4H ₂ O Calcd, %. C. 48.78; H, 4.20; N, 11.98 Found%: C, 48.76; H, 4.34; N, 11.89

Example	R1	R ²	m	Physical properties (Recrystallization solvent)
119	ОН	CI	2	pale brown crystals (CICH ₂ CH ₂ CHeOH) mp.240-245°C (decomposition) Elemental analysis for C ₁₇ H ₁₉ CIN ₄ O-1/2H ₂ O calcd, %: C, 60.09; H, 5.93; N, 16.49 Found%: C, 60.32; H, 5.72; N, 16.41
120	Me	CI	2	pale brown crystals [trifluoroacetate] (EIOH) mp_201-202°C Elemental analysis for Cl ₃ H ₃ ClN ₄ -CF ₂ O ₂ H-5/4H ₂ O Cald %, C, 51, 62; H, 5.31; N, 12.04 Found%: C, 51.82; H, 5.12; N, 12.22
121	CF ₃	CI	2	colorless crystals [trifluoroacetate] ((EIOH) mp,239-235°C Elemental analysis for C1gH1gClFgN_rCF_5CO_H Cackd % C, 48.35; H, 3.85; N, 11.28 Found%: C, 48.31; H, 3.88; N, 11.21
122	Ph	Н	2	colorless crystals [hydrochloride](EtOH) mp,191.5-192.5°C Elemental analysis for C ₂₈ H ₂₄ N ₄ :2HCH+I ₂ O Calcd, %: C, 61.74; H, 6.31; N, 12.52 Found%: C, 61.69; H, 6.51; N, 12.44
123	Ph	CI	3	colorless fine needles[trifluoroacetate] (EICH) mp_260-263°C (decomposition) Elemental analysis for C ₂₄ H ₂₆ CIN ₄ . CF ₂ CO ₂ H Calcd.%. C, 60.17; H, 5.05; N, 10.80 Found%. C, 59.94; H, 5.08; N, 10.80

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(continued)

Example	R2	В	w	Physical properties (Recrystallization solvent)
125	CI	CI	СН	colorless crystals [trifluoroacetate](MeOH) mp.249-255°C (decomposition) Elemental analysis for C ₂₃ H ₂₂ Cl ₂ N ₄ -CF ₅ CO ₂ H Calcd. %: C, 55.67; H, 4.30; N, 10.39 Found%: C, 55.75; H, 4.00; N, 10.47
126	CI	Me	СН	colorless fine needles[trifluoroacetate] ((MoCH) mp.255-262°C (decomposition) Elemental analysis for $C_{24}H_{25}CIN_4$ - CF_5CO_2 H Calcd.%: C, 60.17; H, 5.05; N, 10.80 Found%: C, 59.95; H, 5.03; N, 10.79
127	CI	MeO	СН	pale yellow crystals (EtOH) mp.169-170°C Elemental analysis for C ₂₄ H ₂₅ ClN ₄ O-1/2H ₂ O Calcd.%: C, 67.05; H, 6.10; N, 13.03 Found%: C, 67.32; H, 6.06; N, 13.02
128	CI	н	N	colorless crystals [trifluoroacetate](MeOH) mp;260-268°C (decomposition) Elemental analysis for C ₂₂ H ₂₂ ClN ₅ -CF ₃ CO ₂ H Calcd.%: C, 56.98; H, 4.58; N, 13.84 Found%: C, 56.76; H, 4.47; N, 13.82

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Example	R²	R³	Physical properties (Recrystallization solvent)
129	CI	Ç _I	colorless prisms (MeOH) mp,191–193°C Elemental analysis for C ₂₃ H ₂₃ CiN ₄ Caled.%. C, 70.67; H, 5.93; N, 14.33 Found%: C, 70.70; H, 6.08; N, 14.28
130	CI	HN	colorless crystals (AcOEt) mp,156.5–157.5°C Elemental analysis for C ₂₂ H ₂₃ CIN ₄ Calcd.3: C, 70.67; H, 5.93; N, 14.33 Found8: C, 70.64; H, 5.92; N, 14.21
131	CI	HN O	colorless crystals (EtOH) mp,169-171°C Elemental analysis for C ₂₂ H ₂₁ CIN ₄ O Calcd.X. C, 672.6; H, 5.39; N, 14.26 FoundX: C, 67.31; H, 5.55; N, 14.32
132	CI	H ₂ N	colorless crystals [trifluoroacctate] (iso-PrOH) mp.158-163°C (decomposition) Elemental analysis for C ₃₂ H ₂ CIN ₃ ·2CF ₅ CO ₂ H·3/2H ₂ O Calod.k: C, 49.96; H, 4.42; N, 10.60 Founds: C, 49.94; H, 4.41; N, 10.73
133	Me	H ₂ N N	pale brown crystals (AcOEt) mp,88-89°C Elemental analysis for C ₂₄ H ₂ ,N ₅ ·H ₂ O Calcd.½: C, 71.44; H, 7.24; N, 17.36 Found½: C, 71.25; H, 7.23; N, 17.03

	Example		Physical properties	
5	Lxumpic		(Recrystallization solvent)	
-			colorless fine needles[fumarate](EtOH)	
			mp,261-272°C (decomposition)	
		(\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	Elemental analysis for	
10	134	1 1 1	C22H21CIN4-1/2C4H4O4-5/2H2O	
,,,	134		Calcd.%: C, 60.06; H, 5.88; N, 11.67	
			Found%: C, 60.07; H, 5.89; N, 11.60	
		√ N CI	Specific rotation	
15			[α] _D ²⁰ : -12.0° (c=0.1, DMSO)	
		_	colorless crystals [trifluoroacetate]	
		HŅ BL	(EtOH)	
	135	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	mp,215-221°C (decomposition)	
20		I h	Elemental analysis for	
			C ₂₃ H ₂₇ CIN ₄ • CF ₃ CO ₂ H	
		N CI	Calcd.%: C, 59.00; H, 5.55; N, 11.01	
			Found%: C, 58.85; H, 5.63; N, 11.05	
25		_	pale brown crystals [trifluoroacetate]	
		HN Ph	(MeOH-iso-PrOH)	
		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	mp,225-232℃ (decomposition)	
	136	Ĩ »	Elemental analysis for	
30		\nearrow	C ₂₂ H ₂₅ CIN ₄ · CF ₃ CO ₂ H	
		N CI	Calcd.%: C, 58.24; H, 5.29; N, 11.32	
			Found%: C, 58.09; H, 5.29; N, 11.32	
		^	pale brown crystals [trifluoroacetate]	
35	HN Ph	(EtOH)		
	137	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	mp,224-224.5℃	
			Elemental analysis for	
			C ₂₁ H ₂₁ CIN ₄ S·CF ₃ CO ₂ H·3/2H ₂ O	
40		1	N CI	Calcd.%: C, 51.35; H, 4.68; N, 10.41
			Found%: C, 51.65; H, 4.32; N, 10.16	

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Example	R ¹	Physical properties
		(Recrystallization solvent)
		colorless crystals (AcOEt)
		mp,130-131°C
138	n~Bu	Elemental analysis for C ₂₁ H ₂₇ ClN ₄
		Calcd.%: C, 68.00; H, 7.34; N, 15.10
		Found%: C, 67.76; H, 7.59; N, 14.96
		colorless crystals [trifluoroacetate](EtOH)
	_	mp,139−139.5°C
139		Elemental analysis for
		C ₂₃ H ₂₉ CIN ₄ -3/2CF ₃ CO ₂ H-H ₂ O
		Calcd.%: C, 53.29; H, 5.59; N, 9.56
		Found%: C, 53.23; H, 5.33; N, 9.56
		pale brown crystals (AcOEt-iso-Pr ₂ 0)
i		mp,230-234°C (decomposition)
140	Bn	Elemental analysis for C24H25CIN4 · 1/4H2O
		Calcd.%: C, 70.40; H, 6.28; N, 13.68
		Found%: C, 70.41; H, 6.27; N, 13.54
		pale yellow crystals [methanesulfonate]
		(MeOH)
141		mp,196-207°C (decomposition)
		Elemental analysis for
	/ ~ /	C25H25CIN4-2CH3SO3H-H2O
		Calcd.%: C, 51.71; H, 5.62; N, 8.93
		Found%: C. 51.59: H. 5.42: N. 8.87

Example	R ¹	Physical properties (Recrystallization solvent)
142	Me	colorless crystals [fumarate](MeOH) mp.224-229°C (decomposition) Elemental analysis for C ₂₄ H ₂₅ ClN ₄ · C ₄ H ₄ O ₄ · H ₂ O Calcd, žr. G. 82.39; H. 5.80; N. 10.39 Foundži: O, 62.46; H. 5.51; N. 10.42
143	OMe	colorless crystals [fumarate](EtOH) mp.213.5-216°C (decomposition) Elemental analysis for C ₂₄ H ₂₅ CIN ₄ 0 · C ₄ H ₄ O ₄ · 1/4H ₂ O Caled. %: C, 62.10; H, 5.49; N, 10.35 Found%: C, 61.94; H, 5.45; N, 10.30
144	SMe	colorless crystals [trifluoroacetate] (MeOH-iso-Pr ₂ O) mp_2S3-2S7*C(decomposition) Elemental analysis for C ₂₄ H ₂₆ CIN,S·CF ₂ CO ₂ H-1/2H ₂ O Calcd.S·C, 55.76; H, 4.86; N, 10.00 Found%: C, 55.67; H. 4.55; N, 9.99
145	Me s	colorless crystals [trifluoroacetate](EtOH) mp,218-225°C (decomposition) Elemental analysis for C ₂₄ H ₂₅ CIN ₄ OS-CF ₃ CO ₂ H Calcd &: C, 55.07; H, 4.62; N, 9.88 Found%: C, 54.91; H, 4.69; N, 9.77
146	Ms	coloriess crystals [trifluoroacetate](MeOH) mp.270-277°C (decomposition) Elemental analysis for C ₂₄ H ₂₂ CIN ₂ O ₂ S-CF ₃ CO ₂ H Calcd.S: C, 53.56; H, 4.49; N, 9.61 Found%: C, 53.51; H, 4.50; N, 9.62

		,
Example	R ¹	Physical properties (Recrystallization solvent)
147	F	colorless crystals [fumarts(JEtOH) mp,192–198°C (decomposition) Elemental analysis for C ₂₂ H ₂₂ CIFN ₄ ·C ₄ H ₄ O ₄ ·H ₂ O Calod.%: C, 59.72; H, 5.20; N, 10.32 Found%: C, 59.81; H, 5.07; N, 10.33
148	Ĵ,	colorless crystals [furmarate](MeOH-iso-PrOH) mp,184-187°C (decomposition) Elemental analysis for C ₂₁ H ₂₂ ClFN ₄ ·C ₄ H ₄ O ₄ ·H ₂ O Calcd.S. C, 59.72; H, 5.20; N, 10.32 FoundS: C, 60.00; H, 4.91; N, 10.34
149	∫∫ ^F	coloriess crystals [fumarate](MeOH) mp.204-209°C (decomposition) Elemental analysis for C ₂₃ H ₂₂ ClFN ₄ ·C ₄ H ₄ O ₄ ·H ₂ O Calcd.S. C, 59.72; H, 5.20; N, 10.32 FoundS: C, 59.53; H, 492; N, 10.41
150	F	colorless crystals [trifluoroscetate](EtOH) mp,260-263°C (decomposition) Elemental analysis for Cg ₂ H ₁₂ CiF ₄ N ₄ · CF ₃ CO ₂ H· H ₂ O Calcd.%: C, 50.47; H, 3.73; N, 9.42 Found%: C, 50.33; H, 3.53; N, 9.51
151		coloriess crystals [trifluoroacetate](MeOH) mp,259-261°C (decomposition) Elemental analysis for C ₂ H ₁₂ ClF ₅ N ₄ ·CF ₅ CO ₂ H Calcd.S. C, 50.48; H, 3.22; N, 9.42 FoundS: C, 50.28; H, 3.28; N, 9.48

Example	R1	Physical properties
Example		(Recrystallization solvent)
		coloriess crystals [methanesulfonate]
		(EtOH)
	N/	mp,195-202℃ (decomposition)
152		Elemental analysis for
		C ₂₂ H ₂₂ CIN ₅ • CH ₃ SO ₃ H • 5/4H ₂ O
		Calcd.%: C, 54.11; H, 5.63; N, 13.72
		Found%: C, 54.13; H, 5.45; N, 13.63
		colorless crystals [fumarate](MeOH-EtOH)
		mp,181-185.5℃ (decomposition)
153		Elemental analysis for
100	N	C22H22CIN5 · C4H4O4 · H2O
	,	Calcd.%: C, 59.37; H, 5.37; N, 13.31
		Found%: C, 59.37; H, 5.11; N, 13.37
		pale yellow fine needles [trifluoroacetate]
		(EtOH)
	∕ N	mp,197.5-204°C (decomposition) Elemental analysis for C ₂₂ H ₂₂ ClN ₅ ° CF ₃ CO ₂ H • 1/4H ₂ O
154	ار ا	
	/ //	
		Calcd.%: C, 56.47; H, 4.64; N, 13.72
		Found%: C, 56.45; H, 4.58; N, 13.72
		colorless crystals [trifluoroacetate](EtOH)
	Ph	mp,250-255°C (decomposition)
155		Elemental analysis for C29H27CIN4-CF3CO2H
	/ 🏏	Calcd.%: C, 64.08; H, 4.86; N, 9.64
		Found%: C, 63.81; H, 4.92; N, 9.63
		colorless crystals [trifluoroacetate](EtOH)
	.OPh	mp,144.5-145.5℃
156		Elemental analysis for
		C ₂₉ H ₂₇ ClN ₄ O • CF ₃ CO ₂ H • 3/2H ₂ O
		Calcd.%: C, 59.66; H, 5.01; N, 8.98
- 1		Found%: C. 59.44: H. 4.71: N. 9.04

Example	R¹	Physical properties
		(Recrystallization solvent)
		pale green crystals[trifluoroacetate](EtOH)
	.CF₁	mp,174-175°C
157	(\	Elemental analysis for
		C ₂₄ H ₂₂ CIF ₃ N ₄ ·CF ₃ CO ₂ H·5/4H ₂ O
		Calcd.%: C, 52.44; H, 4.32; N, 9.41
		Found%: C, 52.54; H, 4.19; N, 9.53
		colorless crystals [trifluoroacetate](MeOH)
	^	mp,231-241°C (decomposition)
158	9	Elemental analysis for
)=	C21H21CIN4O-CF3CO2H-1/2H2O
	ı '	Calcd.%: C, 54.82; H, 4.60; N, 11.12
		Found%: C, 54.73; H, 4.42; N, 11.21
	^	colorless crystals [trifluoroacetate](EtOH)
		mp,256-261°C (decomposition)
159	\$ 7	Elemental analysis for
133) /	C21H21CIN4S · CF3CO2H · 1/4H2O
	•	Calcd.%: C, 53.59; H, 4.40; N, 10.87
		Found%: C, 53.53; H, 4.33; N, 10.90
1		colorless crystals [trifluoroacetate](MeOH)
	^	mp,270-273°C (decomposition)
160	HN	Elemental analysis for
)==N	C ₂₀ H ₂₁ CIN ₈ • CF ₃ GO ₂ H • 1/2H ₂ O
	•	Calcd.%: C, 52.44; H, 4.60; N, 16.68
		Found%: C, 52.15; H, 4.74; N, 16.95
		pale brown crystals [trifluoroacetate]
	^	(EtOH-Et ₂ O)
161	s´ 🦻	mp,203-203.5°C
) _ N	Elemental analysis for C ₂₀ H ₂₀ ClN ₅ S·CF ₂ CO ₂ H
	•	Calcd.%: C, 51.61; H, 4.13; N, 13.68
		Found%: C. 51.48: H. 4.22: N. 13.52

Example	mple R ¹	Physical properties
Example		(Recrystallization solvent)
		pale yellow crystals [hydrochloride](iso-PrOH)
	△ .F	mp,245-249℃(decomposition)
162		Elemental analysis for C24H25FN4 · 2HCI · 3/4H2O
		Calcd.%: C, 60.70; H, 6.05; N, 11.80
		Found%: C, 60.81; H, 5.93; N, 11.72
		colorless crystals [hydrochloride](EtOH)
	. [.	NMR spectrum δ
163	Y	(DMSO-d ₆)ppm:1.30-1.40(2H,m),1.55-1.70(1H,m),1.70
103	\\\	-1.80(4H,m),2.65-2.80(2H,m),3.10-3.25(2H,m),3.17(3H
	Į.	,s),4.73(2H,t,J=7.5Hz),7.97(1H,t,J=7.5Hz),8.04(1H,t,J=
		7.5Hz),8.55-8.65(2H,m),8.84(1H,brs),9.06(1H,brs)
		pale brown crystals (AcOEt)
	N	mp,176-177.5℃
164		Elemental analysis for C ₂₃ H ₂₅ N ₅
		Calcd.%: C, 74.36; H, 6.78; N, 18.85
		Found%: C, 74.09; H, 6.90; N, 18.69
		colorless crystals [hydrochloride]
	∠CF ₃	(MeOH-iso-PrOH)
165	Cr3	mp,>300°C
100		Elemental analysis for C ₂₅ H ₂₅ F ₃ N ₄ ·2HCl·1/2H ₂ O
1		Calcd.%: C, 57.70; H, 5.42; N, 10.77
		Found%: C, 57.72; H, 5.12; N, 10.79
		pale yellow crystals (iso-PrOH)
	· 🔊	mp,166−167°C
166		Elemental analysis for C ₂₂ H ₂₄ N ₄ O • H ₂ O
]		Calcd.%: C, 69.82; H, 6.92; N, 14.80
1		Found%: C, 69.53; H, 6.97; N, 14.59

Example	R ^t	Physical properties (Recrystallization solvent)
167	HN	colorless crystals [hydrochloride] (EtOH) mp_218-219°C Elemental analysis for C ₂₁ H ₂₄ N ₆ ·3HCl Calcd.%: C, 53.68; H, 5.79; N, 17.89 Found%: C, 53.63; H, 6.01; N, 17.89
168	5	pale yellow crystals [hydrochloride] (MeOH) mp.293-298°C (decomposition) Elemental analysis for C ₂₁ H ₂₁ N ₃ S-2HOI+H ₂ O Galcd.8: C, 53.84; H, 5.81; N, 14.95 Found8: C, 53.59; H, 5.71; N, 14.82
169	\$	pale yellow crystals [hydrochloride] (EtOH) mp,186–199°C Elemental analysis for C ₂₂ H _{2,N} S-2HCl-3H ₂ O Calod.S: C, 52.48; H, 6.41; N, 11.13 FoundS: C, 52.44; H, 6.68; N, 11.13
170	S Me	pale yellow crystals [trifluoroacetate] (EtOH) mp.228-229°C Elemental analysis for C ₂₃ H ₂₃ N ₃ S-3/2CF ₅ CO ₂ H-1/2H ₂ O Calcd.S. C, 54.73; H, 5.03; N, 9.82 FoundS: C, 54.46; H, 4.91; N, 10.00
171	S Me	pale yellow crystals [hydrochloride] (EtOH) mp.274-277°C (decomposition) Elemental analysis for C ₂₃ H ₂₈ N ₄ S-2HC15/4H ₂ O Calcd.\$C, C, 56.84; H, 6.33; N, 11.53 Found\$C, C, 56.79; H, 6.11; N, 11.51

[Example	R ¹	R²	Physical properties (Recrystallization solvent)
	172	S Me	CI	colorless arystals [trifluoroacetate] (E:OH) mp.189–190°C Elemental analysis for Cp.Hp_CIN,S-3/2CF_3CO,H Calod.3: C, 515-54; H, 424; N, 9.83 Found%: C, 515-54; H, 429; N, 9.85
	173	s Me	GI	colorless crystals [trifluoroacetate] (EtOH) mp.184-195°C Elemental analysis for C ₂₂ H ₃₂ OlN ₃ S-5/40F ₃ CO ₂ H Calcd.3: C, S3.16: H, 4.42; N, 10.39 Found's: C, S3.16: H, 4.39: N, 10.39
	174	HN	Мо	pale brown crystals [hydrochloride] (ECOH) mp_245.5-246.5°C Elemental analysis for C ₂₅ H ₂₆ N ₆ -2HCI-3/2H ₂ O Calcd.3: C, 57.52; H, 6.58; N, 15.24 Found%: C, 57.55; H, 6.33; N, 15.23
	175	MeN	Ме	pale brown orystals [hydrochloride] (EtOH) mp.224-225°C Elemental analysis for C ₂₂ H ₂₂ N ₃ -2HOI-5/2H ₂ O Calcd.k: G, 562.H, 507; N, 14.25 Founds: G, 55.95; H, 5.70; N, 14.23
	176	н	, F	coloriess prisms[trifluoroacetate] (EEOH-iso-Pr ₂ O) mp.189.5–192.5°C Elemental analysis for C ₂₂ H ₂₂ FN ₄ O-CF ₂ CO ₂ H Calcd.8: C, 59.52: H, 4.80: N, 11.11 Found'S: C, 59.41: H, 4.89: N, 11.16

Example	R²	Physical properties
		(Recrystallization solvent)
1		colorless crystals [trifluoroacetate]
i		(EtOH)
		mp,214.5-215.5℃
177	OPh	Elemental analysis for
1		C29H28N4O-CF3CO2H-1/2H2O
		Calcd.%: C, 65.14; H, 5.29; N, 9.80
		Found%: C, 65.40; H, 5.07; N, 9.85
		colorless crystals (MeOH-iso-PrOH)
1		mp,191~194℃
178	NHPh	Elemental analysis for C ₂₉ H ₂₉ N ₅
		Caled.%: C, 77.82; H, 6.53; N, 15.65
		Found%: C, 77.76; H, 6.59; N, 15.56
		pale yellow crystals [hydrochloride]
-	NHMe	(iso-PrOH)
1		mp,209-210°C
179		Elemental analysis for
		C24H27N5 · 2HCI · 7/4H2O
		Calcd.%: C, 58.83; H, 6.69; N, 14.29
		Found%: C, 58.88; H, 6.51; N, 14.13
		colorless crystals [hydrochloride]
1		(MeOH)
		mp,205-206.5°C
180	NMe ₂	Elemental analysis for
		C25H29N5 · 2HCI · 5/2H2O
1		Calcd.%: C, 58.02; H, 7.01; N, 13.53
		Found%: C, 58.01; H, 7.02; N, 13.50
		coloriess crystals [hydrochloride]
		(EtOH)
	l	mp,210-212°C
181	N ~	Elemental analysis for
	#	C ₂₆ H ₂₉ N ₅ · 2HCI · H ₂ O
		Calcd.%: C, 62.15; H, 6.62; N, 13.94
1		Found%: C, 61.99; H, 6.44; N, 13.85

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Example	R²	Physical properties
		(Recrystallization solvent)
		colorless crystals [hydrochloride]
		(iso-PrOH)
Į.		mp,244-245°C
182	NHBn	Elemental analysis for
Í		C ₃₀ H ₃₁ N ₅ -2HGI-3/4H ₂ O
		Calcd.X: C, 65.75; H, 6.35; N, 12.78
		Found%: C, 65.81; H, 6.13; N, 12.68
		pale yellow crystals [hydrochloride]
		(EtOH)
	\n\\\	mp,190−193°C
183	# []	Elemental analysis for
	~n	C ₂₉ H ₃₀ N ₆ ·3HCI·2H ₂ O
		Calcd.%: C, 57.29; H, 6.13; N, 13.82
		Found%: C, 57.46; H, 5.98; N, 13.77
		pale yellow crystals [hydrochloride]
	N NMe	(EtOH)
		mp,231.5-232°C
184		Elemental analysis for
į.		C28H34N6-3HCI-3/4H2O
		Calod.%: C, 58.23; H, 6.72; N, 14.55
		Found%: C, 58.12; H, 6.93; N, 14.46
l		colorless needles [hydrochloride]
I		(EtOH)
-	`N ^	mp,187-189℃
185	ŢJ	Elemental analysis for
	~	C ₂₈ H ₃₃ N ₅ • 2HCl • 3/4H ₂ O
į		Calcd.%: C, 63.93; H, 6.99; N, 13.31
		Found%: C, 64.05; H, 6.93; N, 13.32
i		colorless crystals [hydrochloride]
- 1	`\;`	(EtOH-iso-PrOH)
1		mp,194~195°C
186		Elemental analysis for
ŀ		C ₂₇ H ₃₁ N ₅ O • 2HCl • 3/2H ₂ O
1		Calcd.%: C, 59.89; H, 6.70; N, 12.93
- 1		Found%: C, 59.72; H, 6.64; N, 12.85

Example 187

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1-[2-(N-n-Butyl-4-piperidyl)ethyl]-4-chloro-1H-imidazo[4,5-c]quinoline hydrochloride

5 [0119] To a suspension of 1.20 g of 4-chloro-1-[2-(4-piperidyl)ethyl]-IH-imidazo-[4,5-c]quinoline trifluoroacetate and 0.77 g of potassium carbonate in 6 ml of N.N-dimethylformamide, 0.30 ml of n-butyl bromide was added dropwise at room temperature, and the mixture was stirred for 5 hours. The reaction mixture was adjusted to pH 10 with 10% aqueous sodium hydroxide solution, and extracted with ethyl acetate. The extract was washed successively with water

and saturated brine, and dried, and then the solvent was evaporated to give 0.92 g of a pale brown liquid. The resulting liquid was dissolved in letrahydrofuran. The solution was filtered on silica gel, and the filtrate was concentrated to give 0.87 g of a coloriess solid. Hydrochloride was prepared in a conventional method. Recrystallization from a mixture of methanol and ethyl acetate gave coloriess crystals having the mething point of from 1.44 to 158°C.

Elemental analysis for C ₂₁ H ₂₇ ClN ₄ - 2HCl - 1/2H ₂ O							
Calculated % C, 55.70; H, 6.68; N, 12.37							
Found %	C, 55.80;	H, 6.65;	N, 12.44				

Example 188

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1-[2-(N-Acetyl-4-piperidyl)ethyl]-4-chloro-1H-imidazo[4,5-c]quinoline

[0120] To a solution of 0.80 g of 4-chbror-1/2-(4-piperidy)ethyl)-11-t-midszo-/4,5-c)quinoline trifluoroacetate in 4 mi of pyridine, 2 mi of acetic anhydride was added, and the mixture was stirred at room temperature for 1 hour. After the reaction, the solvent was evaporated. The residue was added with isopropanol and disopropy ether, and the precipitated crystals were collected by filtration, and washed with disopropyl ether to give 0.45 g of colorless crystals. Recrystalization from a mixture of methylene chloride and disopropyl ether gave colorless crystals having the meiting point of from 183 to 186.5°C.

Elemental analysis for C ₁₉ H ₂₁ ClN ₄ O							
Calculated % Found %	C, 63.95;	H, 5.93;	N, 15.70				
Found %	C, 63.81;	H, 5.87;	N, 15.61				

[0121] In accordance with the methods of Examples 187 and 188, the compounds of Examples 189 through 194 were obtained.

	Example	R1	В	R ³	m	Physical properties
						(Recrystallization solvent)
5			1		l	colorless crystals (iso-PrOH)
				MeN ^		mp,167-168℃
	189	Ph	Н	men	2	Elemental analysis for C24H25CIN4
						Calcd.%: C, 71.19; H, 6.22; N, 13.84
10					L	Found%: C, 71.00; H, 6.18; N, 13.56
						coloriess crystals [hydrochloride]
						(EtOH)
				BnN	2	mp,235-246°C (decomposition)
15	190	Н	CI			Elemental analysis for
						C24H24Cl2N4+HCl+1/4H2O
						Calcd.%: C, 60.01; H, 5.35; N, 11.66
						Found%: C, 60.01; H, 5.62; N, 11.67
20						colorless crystals [hydrochloride]
						(EtOH)
				BnN		mp,248-257°C (decomposition)
	191	н	н	5111	1	Elemental analysis for
25				\sim		C ₂₃ H ₂₃ CIN ₄ +HCI-1/4H ₂ O
						Calcd.%: C, 63.96; H, 5.72; N, 12.97
						Found%: C, 63.98; H, 5.80; N, 12.93
						colorless crystals (CH ₂ Cl ₂ -iso-Pr ₂ O)
30						mp,154.5-160°C
	192	Ph	h H	AcN	2	Elemental analysis for
						C25H25CIN4O · 1/8H2O
				,		Calcd.%: C, 69.00; H, 5.85; N, 12.87
35						Found%: C, 68.78; H, 5.78; N, 12.71

Example	R³	m	Physical properties (Recrystallization solvent)
193	BnN	1	colorless crystals [hydrochloride] (MaOH-ise-Pr ₂ O) mp_289-280°C (decomposition) Elemental analysis for C ₂₂ H ₁₄ N ₄ ·2HCl·3/4H ₂ O Calod.: C, 62.37; H, 6.26; N, 12.65 Found%: C, 62.36; H, 6.45; N, 12.60
194	BnN	2	colorless crystals [hydrochloride] (MaOH-iso-Pr ₂ O) mp.150-156°C (decomposition) Elemental analysis for C ₁₄ H ₂₈ N ₁ -2Hcl-1/2H ₂ O Calcd. C, 63.71, B, 646; N, 12.38 Found's: C, 63.90; H, 6.88; N, 12.11

Example 195

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4-Chloro-1-[2-[N-(4-fluorophenylsulfonyl)-4-piperidyl]ethyl]-1H-imidazo-[4,5-c]quinoline

[0122] To a suspension of 0.50 g of 4-shoro-1-(2-(4-p)peridy)lethyl-1H-indiazo-(4,5-e)quinoline trifluoroacetale and 0.32 g of polassium carbonate in 2 ml of N N-dimethylformamide, a solution of 0.23 g of polisoroacensesullonyl, choice in 3 ml of N N-dimethylformamide was added dropwise at room temperature, and the mixture was stirred for 5 hours. The reaction mixture was adjusted to p+10 of with 0% aqueous sodium hydroxide solution, and extracted with ethyl acetate. The extract was washed successively with water and saturated brine, and dried, and then the solvent was evaporated to give 0.35 g of a colorless solid. Recrystallization from a mixture of methanol, ethanol and water gave colorless crystals having the metiting point of from 175 to 178.5°C.

Elemental analysis for C ₂₃ H ₂₂ CIFN ₄ O ₂ S							
Calculated % C, 58.41; H, 4.69; N, 11.85 Found % C, 58.43; H, 4.52; N, 11.88							
Found %	C, 58.43;	H, 4.52;	N, 11.88				

Example 196

1-[2-(N-Methanesulfonyl-4-piperidyl)ethyl]-4-phenoxy-1H-imidazo[4,5-c]-quinoline

[0123] To a solution of 1,00 g of 4-phenoxy-1-[2-(4-piperdy)Pethyl-H-imidazo-14,5-cipunionine trilluroracetate and 0,57 ml of triethylamine in 10 ml of methylene chloride, 0.16 ml of methanesulfonyl chloride was added dropwise at room temperature, and the mixture was stirred for 1.5 hours. The reaction mixture was added with water, and extracted with methylene chloride. The extract was washed with water, and dried, and then the solvent was evaporated to give a colorises liquid. The resulting colorises liquid was solidified with othyl accetate, and the solid was washed with diethyl either to give 0.80 g of coloriess crystals. Recrystalitization from a mixture of methylene chloride and ethyl acetate gave coloriess crystals having the melting point of from 17.35 to 176°C.

Elemental analysis for C ₂₄ H ₂₆ N ₄ O ₃ S						
Calculated % C, 63.98; H, 5.82; N, 12.44						
Found %	C, 64.01;	H, 5.96;	N, 12.28			

[0124] In accordance with the method of Example 196, the compounds of Examples 197 through 199 were obtained.

	Example	R ^A	Physical properties (Recrystallization solvent)
15	197	Ts	colorless crystals (AcOEI-iso-Pr ₂ O) mp.201.5-202**C Elemental analysis for C ₃₀ H ₃₀ N ₄ O ₂ S Calcd.**C, 68.42: H, 5.74; N, 10.64 Found**C, 68.46: H, 5.83; N, 10.55
20	198	EtO ₂ C	colorless crystals (AcOEt-iso-Pr ₂ O) mp.132-133*C Elemental analysis for C ₂₆ H ₂₆ N ₄ O ₃ Catcd.%: C, 70.25; H, 6.35; N, 12.60 Found%: C, 70.15; H, 6.34; N, 12.60
25	199	BnO ₂ C	yellow liquid NMR apectrum δ (CDCl ₂)ppm: 1.31 (2H,brs),1.50-1.70(1H,m),1.78(2H,brs),2.00(2H,q,J=7.5Hz),2.81(2H,brs),4.23(2H,brs),4.83(2H,L ₃ -7.5Hz),5.13(2H,brs),4.25(2H,L ₃ -7.5Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.39(2H,d,J=7Hz),7.30-7.40(6H,m),7.30(2H,d,J=7Hz),7.30(2H,d,
30			7.44(2H,Ju-7H)2,7.50(1H,dd,J=8.5,1H2),7.57(1H,t d,J=8.5,1H2),7.90(1H,dd,J=8.5,1H2), 7.94(1H,s),8.04(1H, dd,J=8.5,1Hz) IR spectrum v(IIq.) cm*1:1698 Mass spectrum m/z:506(M*)

Example 200

4-[2 - (4-Amino-1H-imidazo [4,5-c]quinolin-1-yl)ethyl]-N-methyl-1-piperidine-carbothioamide

[0125] A suspension of 0.50 g of 4-amino-1-[2-(4-piperidyl)ethyl]-1H-imidazo[4,5-c]-quinoline and 0.37 g of methyl-isothicoyanate in 10 ml of methylene chloride was stirred at room temperature for 1 hour, and then the precipitated crystals were collected by filtration to give 0.56 g of coloriess crystals. Recrystallization from a mixture of methylene chloride and methanol gave coloriess crystals having the melting point of from 216 to 218°C.

Elemental analysis for C ₁₉ H ₂₄ N ₆ S · 1/2H ₂ O						
Calculated % C, 60.45; H, 6.67; N, 22.26						
Found % C, 60.79; H, 6.66; N, 21.97						

[0126] In accordance with the method of Example 200, the compound of Example 201 was obtained.

Example 201

4-[2-(4-Chloro-2-phenyl-1H-imidazo[4,5-c]quinolin-1-yl)ethyl]-N-methyl-1-piperidinecarbothioamide

[0127]

[-..

Appearance: colorless crystals Recrystallization solvent: methanol mp: 215-220°C (decomposition)

Elemental analysis for C ₂₅ H ₂₆ CIN ₅ S						
Calculated % C, 64.71; H, 5.65; N, 15.09						
Found %	C, 64.80;	H, 5.62;	N, 14.96			

Example 202

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1-[2 -(1-Amidino-4-piperidyl)ethyl]-4-chloro-2-phenyl-1H-imidazo[4,5-c]-quinoline hydrochloride

[0128] A solution of 0.75 g of 4-chloro-2-phenyl-1-[2-(4-piperidyl)ethyl]-IH-imidazo-[4,5-c]quinoline, 0.40 g of 1Hpyrazole-1-carboxyamidine hydrochlorido and 0.39 ml of interhylamine in 5 ml of IN-N-dimetrylformamide was dated at room temperature for 19 hours. The reaction solution was concentrated and the residue was added with ethanol, and then the precipitated crystals were collected by filtration to give 0.51 g of colorless crystals. Recrystallization from ethanol gave colorless crystals having the method report of the 200 to 279°C (decomposition).

Elemental analysis for C ₂₄ H ₂₅ ClN ₆ · HCl · 1/2H ₂ O					
Calculated %	C, 60.25;	H, 5.69;	N, 17.57		
Found %	C, 60.47;	H, 5.61;	N, 17.36		

[0129] As an example of the excellent effects of the compounds according to the present invention, experimental results of inhibitory actions against production of TNF- α and IL-1 β in human cells will be shown below.

1. Preparation of blood cells for culture

[0130] About 50 mL of whole blood was collected from adult healthy volunteers by venepuncture into a plastic tube which containing 170 µL of Novo-heparin 1000 (Novo-hovdisk A/S). Then, PBMCs (Perhiperal Blood Mononucleat Cells) were prepared using a cell separation tube, Leuco-PREP™ (Becton Dickinson), and cultured with RPMI-160 medium (Nissui Pharmaceutical Co.) containing 2 mM L -glutamine (Life Technologies), 2.5 Uml penicillin-2.5 µg/mL streptomycin solution (Life Technologies) supplemented with 10% fetal call serum (Intergen Company) at 1x10⁶ cels/ mL.

Preparation of test compounds

[0131] Test compounds were dissolved in distilled ultra-pure water, dimethyl sulfoxide, or 0.1 N hydrochloric acid at 20 μM, and then sequentially diluted with saline and used. The compounds were examined at concentrations ranging from 10 ¹⁰ M to 10 ⁵ M.

40 3. Treatment of cells with medicaments

[0132] 10 µL of 1 µg/mL lipopolysaccharide (LPS) was added to a 96-well (flat bottom) plate for cell culture, MicroTest III ™ tissue culture plate (Becton Dickinson), containing 180 µL of the PBMCs in the aforementioned medium. After 30 minutes, 10 µL of the solution of the test compound or the solvent was further added to each well, and the plate was covered with a plastic lid and incubated at 37°C for 16 hours in an atmosphere of 5% CO₂.

4. Determination of human TNF-a and human IL-1β

[0133] An enzyme immunoassay by the sandwich method was performed to determine the human TIMF- α and human Li-1 β in the culture supernatant. The anti-toptions enablody (the first-antibody) was diluted and placed in a 98-well microtiter plates for ceating. After the wells were washed, the culture supernatant was appropriately diluted, and then added to each well and incubated. Then the second-antibody against cytokine and the third-antibody against the second-antibody were successively added while applying washing processes between the operations. After the final washing process, a tetrametriy/benz/dine solution (DAKO) was added to each well to start the coloring reaction. The coloring reaction was quenched with 1 N sulfuris acid, and then the absorbance at 450 nm of each well was measured by a microplate reader, M-YmaxTM (Molecular Devices). The concentrations of the cytokines were determined by quantification software, SoftmaxTM (Molecular Devices), in comparison with the calibration curves obtained by using the re-

combinant cytokines as the standards. For determination of human TNF- α , monoclonal anti-human TNF- α (ENDOGEN), polydonal rabbit anti-human TNF- α (Pharma Biotechnologie Hannover), peroxidase conjugated donkey anti-rabbit (gG (Jackson ImmunoHes. Labs.), and recombinant human TNF- α (INTERGEN Company) were used for the first, second- and third-antibodys and the standard for the calibration curve, respectively. For determination of human IL-18 (Geseption) anti-human IL-19 (Gespenseis), HFP conjugated donkey anti-joat (gG (Chemicon International), and recombinant human IL-19 (R&D Systems) were used for the first, second- and third-antibodys and the standard for the calibration curve, respectively.

[0134] In both cases for TNF- α and IL-1 β , the activities of each test compound are shown as percentages (%) of the amount of the cytokine induced by treatment with LPS together with the test compound against the amount of the cytokine induced by treatment solely with LPS.

[0135] Results are shown in tables 1 and 2.

Table 1:

Inhibitory action against TNF- α production in human cells					
Compounds	Administered concentration (µmol/L)				
	0.001	0.01	0.10	1.0	10
Example 89	91	86	90	84	17
Example 110	80	77	26	1	0
Example 113	68	81	86	69	29
Example 117	117	77	71	24	0
Example 118	79	91	88	51	3
Example 121	81	91	49	0	0

Table 2:

Inhibitory action against IL-1 β production in human cells					
Compounds	Administered concentration (µmol/L)				
	0.001	0.01	0.10	1.0	10
Example 89	112	102	96	63	0
Example 110	119	105	85	64	14
Example 113	104	109	116	96	30
Example 117	119	106	111	72	8
Example 118	96	106	102	59	0
Example 121	102	108	87	24	0

[0136] These results clearly indicate that the compounds of the present invention have excellent inhibitory actions against production of TNF and IL-1.

Industrial Applicability

[0137] The compounds of the present invention have excellent inhibitory actions against production of TNF or IL-1 and are extreamely useful as preventive or therapeutic agents of diseases mediated by these cytokines.

Claims

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1. A 1H-imidazopyridine derivative represented by the following general formula or a salt thereof:

- wherein B¹ represents hydrogen atom, hydroxyl group, an alkyl group which may have one or more substituents, a cycloalkyl group which may be substituted, a styryl group which may be substituted, or an anyl group which may have one or more substitutents; R² represents hydrogen atom, an alkyl group, a halogen atom, hydroxyl group, an amino group which may have one or two substituents, a cyclic amino group which may be substituted; ring A represents a homocyclic or a heterocyclic ring which mey be substituted; ring A represents a homocyclic ring which may be substituted and m represents an integer of from 0 to 5; provided when R³ represents unsubstituted piperidino group, at least one of R¹ and R² is not hydrogen atoms.
 - 2. A 1H-imidazopyridine derivative represented by the following general formula or a salt thereof:

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wherein R1 represents hydrogen atom, hydroxyl group, an alloyl group which may have one or more substituents, a cycloalityl group which may be substituted, a styryl group which may be substituted, or an anyl group, which may have one or more substituents, R2 represents hydrogen atom, an alkyl group, a hadiogen atom, hydroxyl group, an amino group which may be substituted, or a phenoxy group which may be substituted, irright and presents a homeocyclic or heterocycle indig which may be substituted with one or more alkyl groups, alkoxyl groups, or halogen atoms; in represents an integer of from 0 to 3; R1 represents hydrogen atom, an alkyl group, bencyl group, the phymethyl group, an laknoyl group which may be substituted, an alknoycarbonyl group, benzyloxycarbonyl group, a thiocarbamoyl group group which may be substituted, an alknoycarbonyl group which may be substituted, an alknoycarbonyl group which may be substituted, an alknoycarbonyl group which may be substituted, an alknoycarbonyl group which may be substituted, an alknoycarbonyl group which may be substituted, an alknoycarbonyl group which may be substituted, or amining group; Yrepresents methylene group, oxygen atom, suffur atom, nitrogen atom, a group represented by NH, or a single bond; and in represents an integer of from 0 to 2.

- The compound or the salt thereof according to claim 1 or claim 2, wherein the ring A is benzene ring or thiophene ring.
- A medicament which comprises as an active ingredient the 1H-imidazopyridine derivative or a pharmacologically acceptable salt thereof according to claim 1 or claim 2.
- The medicament according to claim 4 which is used for preventive or therapeutic treatment of a disease in which a cytokine is mediated.

INTERNATIONAL SEARCH REPORT

A61K31/47

C. DOCUMENTS CONSIDERED TO BE RELEVANT

International application No. PCT/JP99/04381

1-5

1-5

A. CLASSIFICATION OF SUBJECT MATTER Int. C1 C07D471/04, C07D471/14, C07D491/113, C07D495/14, A61K31/435,

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

J. Interferon Res. (1994), 14, P. 81-85

RP. 459505. A (Kyowa Hakko Kogyo Co., Ltd.).

Minimum documentation searched (classification system followed by classification symbols)

Int. C1⁶ C07D471/04, C07D471/14, C07D491/113, C07D495/14, A61K31/435, Int. Cl A61K31/47

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) CAPLUS, REGISTRY (STN)

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×	Further documents are listed in the continuation of Box C.		See patent family annex.
.w.	Special categories of cited documents: document defining the general state of the art which is not considered to be of particular relevance	*T*	later document published after the international filing date or priority date and not in conflict with the application but cited to
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Date of mailing of the international search report

16 November, 1999 (16.11.99)

Japanese Patent Office Form PCT/ISA/210 (second sheet) (July 1992)

Name and mailing address of the ISAV

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Date of the actual completion of the international search

08 November, 1999 (08,11,99)

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Telephone No.

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